



# **STIC Search Report**

## **Biotech-Chem Library**

**STIC Database Tracking Number: 107598**

**TO: Celine Qian**  
**Location: CM1/11C01&11E12**  
**Art Unit: 1636**  
**Wednesday, November 12, 2003**

**Case Serial Number: 09/939209**

**From: Edward Hart**  
**Location: Biotech-Chem Library**  
**CM1-6B02**  
**Phone: 305-9203**

**edward.hart@uspto.gov**

### **Search Notes**

Examiner Qian,

Here are the results of the search you requested.

Please feel free to contact me if you have any questions.

Edward Hart

GenCore version 5.1.6  
Copyright (c) 1993 - 2003 CompuGen Ltd.

OM nucleic - nucleic search, using sw model  
Run on: November 7, 2003, 01:54:16 ; Search time 3916.96 Seconds  
(without alignments)  
10444.222 Million cell updates/sec

Title: US-09-939-209A-3\_COPY\_1\_1000  
Perfect score: 1000  
Sequence: 1 agtcaagaccgctgagc.....gtatgtggcagagcatggt 1000

Scoring table: IDENTITY NUC  
Gapop 10.0 , Gapext 1.0

Searched: 2888711 seqs, 20454813386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database : GenEmbl.\*

- 1: gb.ba.\*
- 2: gb.htg.\*
- 3: gb.in.\*
- 4: gb.om.\*
- 5: gb.ov.\*
- 6: gb.pat.\*
- 7: gb.ph.\*
- 8: gb.pl.\*
- 9: gb.pr.\*
- 10: gb.ro.\*
- 11: gb.sts.\*
- 12: gb.sy.\*
- 13: gb.un.\*
- 14: gb.vi.\*
- 15: em.ba.\*
- 16: em.fun.\*
- 17: em.hum.\*
- 18: em.in.\*
- 19: em.mu.\*
- 20: em.om.\*
- 21: em.or.\*
- 22: em.ov.\*
- 23: em.pat.\*
- 24: em.ph.\*
- 25: em.pl.\*
- 26: em.ro.\*
- 27: em.sts.\*
- 28: em.un.\*
- 29: em.vi.\*
- 30: em.htg.hum.\*
- 31: em.htg.inv.\*
- 32: em.htg.other.\*
- 33: em.htg.mus.\*
- 34: em.htg.pln.\*
- 35: em.htg.rtd.\*
- 36: em.htg.vrt.\*
- 37: em.htg.vrt.\*
- 38: em.sy.\*
- 39: em.htgo.hum.\*
- 40: em.htgo.mus.\*
- 41: em.htgo.other.\*

Pred. No. is the number of results predicted by chance to have a

score greater than or equal to the score of the result being printed,  
and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match %	Length	DB	ID	Description
1	1000	100.0	20300	6	AX451337	AX451337 Sequence
2	1000	100.0	165329	9	AL583850	AL583850 Human DNA
3	617.4	61.7	191699	2	AC031977	AC031977 Homo sapi
4	411.2	41.1	203216	2	AC097328	AC097328 Pan trogl
5	405.2	40.5	181946	2	AC097330	AC097330 Pan trogl
6	402.2	40.2	109081	9	AL358074	AL358074 Human DNA
7	396.8	39.7	164844	9	AC090508	AC090508 Homo sapi
8	395.2	39.5	130349	9	AC011593	AC011593 Homo sapi
9	394.8	39.5	196413	2	AC097269	AC097269 Pan trogl
10	390.8	39.1	189662	9	AC015468	AC015468 Homo sapi
11	387.4	38.7	125685	9	AC073310	AC073310 Homo sapi
12	385.8	38.6	139118	9	AL133477	AL133477 Human DNA
13	382.6	38.3	216021	9	HUAC004787	HUAC004787
14	379.8	38.0	92564	9	AY007685	AY007685 Homo sapi
15	379.4	37.9	161252	2	AC114955	AC114955 Homo sapi
16	378	37.8	202305	9	AC114291	AC114291 Homo sapi
17	373.6	37.4	192139	9	AC021955	AC021955 Homo sapi
18	372	37.2	99596	9	AC106873	AC106873 Homo sapi
19	371.4	37.1	144383	2	EX324168	EX324168 Homo sapi
20	370.4	37.0	252512	2	AC097326	AC097326 Pan trogl
21	368.2	36.8	170646	2	AC123545	AC123545 Pan trogl
22	367	36.7	112762	9	HS323B6	HS323B6 Human DNA s
23	365.4	36.5	163976	9	AC108483	AC108483 Homo sapi
24	365.4	36.5	163998	9	AC022166	AC022166 Homo sapi
25	364.2	36.4	153201	2	AC092863	AC092863 Homo sapi
26	363.6	36.4	151549	9	AC092694	AC092694 Homo sapi
27	357.4	35.7	217985	9	AC021193	AC021193 Homo sapi
28	356.6	35.7	121600	9	HS141H5	HS141H5 Human DNA
29	355.2	35.5	149969	2	AC024180	AC024180 Homo sapi
30	355.2	35.5	162996	9	AC006441	AC006441 Homo sapi
31	354.8	35.5	182430	9	AL590381	AL590381 Human DNA
32	353.6	35.4	164994	2	AC141288	AC141288 Homo sapi
33	352.4	35.2	178254	9	AL731541	AL731541 Human DNA
34	349.6	35.0	50812	9	AL513530	AL513530 Human DNA
35	349.6	35.0	81696	9	AF438327_3	Continuation (4 of
36	349.6	35.0	110000	9	AF438327_2	Continuation (3 of
37	349.2	34.9	171427	9	AC021590	AC021590 Homo sapi
38	348.8	34.9	236551	2	AC094041	AC094041 Rattus no
39	348.6	34.9	134084	9	AC004966	AC004966 Homo sapi
40	348.6	34.9	159817	2	EX322798	EX322798 Danio rer
41	348.4	34.8	247615	2	AC098061	AC098061 Rattus no
42	347.8	34.8	220821	2	AC133068	AC133068 Danio rer
43	347.4	34.7	203216	2	AC097328	AC097328 Pan trogl
44	347.2	34.7	174447	2	AC142552	AC142552 Danio rer
45	347	34.7	137325	2	AC142088	AC142088 Danio rer

ALIGNMENTS

RESULT 1	AX451337	Sequence 3 from Patent WO216653.	20300 bp	DNA	linear	PAT 03-JUL-2002
LOCUS	AX451337	Sequence 3 from Patent WO216653.				
DEFINITION	AX451337					
ACCESSION	AX451337					
VERSION	AX451337.1	GI:21698388				
KEYWORDS		synthetic construct				
SOURCE		artificial sequences.				
ORGANISM		1				
REFERENCE		Levitt,P.R., Mirnice,K., Kodavali,V.C. and Nimgaonkar,V.L.				
AUTHORS		Methods and systems for facilitating the diagnosis and treatment of				
TITLE		schizophrenia				
JOURNAL		Patent: WO 0216653-A 3 28-FEB-2002;				

University of Pittsburgh (US)	
FEATURES	Location/Qualifiers
source	1..20300
	/organism="synthetic construct"
	/mol_type="genomic DNA"
	/db_xref="taxon:32630"
	/note="A genomic sequence containing RGS4 nucleic acid sequence and sequences upstream and downstream to the RGS4 nucleic acid sequence"
BASE COUNT	6157 a 4102 c 3775 g 6266 t
ORIGIN	
Query Match	100.0%; Score 1000; DB 6; Length 20300;
Best Local Similarity	100.0%; Pred. No. 1.2e-185;
Matches 1000; Conservative	0; Mismatches 0; Indels 0; Gaps 0;
QY	1 AGTTCAAGACCGCTGACCAATGTTGAAACCCCATCTCTACTAAATAACAAATTA 60
Db	1 AGTTCAAGACCGCTGACCAATGTTGAAACCCCATCTCTACTAAATAACAAATTA 60
QY	61 GACAGGCATGGTGATACACGCTGTAAATCCAGCTACTTTCGAGGCGGAGGAGAAAT 120
Db	61 GACAGGCATGGTGATACACGCTGTAAATCCAGCTACTTTCGAGGCGGAGGAGAAAT 120
QY	121 CACTTGAACCTCTCGGGGTGAGGTTCGGGGAGCAAGATCATGCCATTGCATCCAGC 180
Db	121 CACTTGAACCTCTCGGGGTGAGGTTCGGGGAGCAAGATCATGCCATTGCATCCAGC 180
QY	181 CCAGGCAACAGAGCGAAATGTCTCAGAAAAAAGCGATTTTATATATATA 240
Db	181 CCAGGCAACAGAGCGAAATGTCTCAGAAAAAAGCGATTTTATATATATA 240
QY	241 TATATATATATACACACACACATATATATATACACATATATATACATATATACA 300
Db	241 TATATATATATACACACACACATATATATATATACACATATATATACATATACA 300
QY	301 TATATACATATATACATATATATATACACATATGTACACATATATATACATA 360
Db	301 TATATACATATATACATATATATATACACATATGTACACATATATATACATA 360
QY	361 TGTATACATATATACATATATATATACACATATATATACATATATATATATA 420
Db	361 TGTATACATATATACATATATATATACACATATATATACATATATATATATA 420
QY	421 CACATATATACATATATATATACACATATATATACATATATATATATATACACA 480
Db	421 CACATATATACATATATATATACACATATATATACATATATATATATATACACA 480
QY	481 TATATATATATACACATATATATATACATATATATACACATATATATATATAC 540
Db	481 TATATATATATACACATATATATATACATATATATATATATATATATATATAC 540
QY	541 ACATATATATACATATATATATACATATATATATACATATATATATATATATAC 600
Db	541 ACATATATATACATATATATATACATATATATATATATATATATATATATATAC 600
QY	601 ACATAC 660
Db	601 ACATAC 660
QY	661 ACACAT 720
Db	661 ACACAT 720
QY	721 GCTCCAGAGTTTCAAGAGGTAGAGTTCATTAACCTCGGGATAGAGGAAAGAGATT 780
Db	721 GCTCCAGAGTTTCAAGAGGTAGAGTTCATTAACCTCGGGATAGAGGAAAGAGATT 780
QY	781 TGACAGCAGTGTATTCTGAGAGGACATTTTCAGTTGATGGCAATAGTAGGGGAATAC 840
Db	781 TGACAGCAGTGTATTCTGAGAGGACATTTTCAGTTGATGGCAATAGTAGGGGAATAC 840
QY	841 ATAAATGTCTAATAAAACCTACTCTGTAGGTAGTTTAAAGAGGTAAACACTATATATA 900
Db	841 ATAAATGTCTAATAAAACCTACTCTGTAGGTAGTTTAAAGAGGTAAACACTATATATA 900
QY	901 TAGTGAAGACCTGTAAACCTAAAGATGGCCAGGATTTAAATGTTTATAGAAGATGG 960
Db	901 TAGTGAAGACCTGTAAACCTAAAGATGGCCAGGATTTAAATGTTTATAGAAGATGG 960
QY	961 CTAAGATGCCAAGCTCAGTGTATGTGGCAGAGGCATGGT 1000
Db	961 CTAAGATGCCAAGCTCAGTGTATGTGGCAGAGGCATGGT 1000
RESULT 2	
AL583850	165329 bp DNA linear PRI 15-NOV-2001
LOCUS	Human DNA sequence from clone Rp11-430G6 on chromosome 1, complete
DEFINITION	sequence.
ACCESSION	AL583850
VERSION	AL583850.5 GI:16973044
KEYWORDS	HTG.
SOURCE	Homo sapiens (human)
ORGANISM	Homo sapiens
REFERENCE	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
AUTHORS	Tracey, A.
TITLE	Direct Submission
JOURNAL	Submitted (15-NOV-2001) Wellcome Trust Sanger Institute, Hinxton, Cambridgeshire, CB10 1SA, UK. E-mail enquiries: humbry@sanger.ac.uk Clone requests: clonerequest@sanger.ac.uk
COMMENT	On Nov 16, 2001 this sequence version replaced gi:15020514. During sequence assembly data is compared from overlapping clones. Where differences are found these are annotated as variations together with a note of the overlapping clone name. Note that the variation annotation may not be found in the sequence submission corresponding to the overlapping clone, as we submit sequences with only a small overlap as described above. This sequence was finished as follows unless otherwise noted: all regions were either double-stranded or sequenced with an alternate chemistry or covered by high quality data (i.e., phred quality >= 30); an attempt was made to resolve all sequencing problems, such as compressions and repeats; all regions were covered by at least one plasmid subclone or more than one M13 subclone; and the assembly was confirmed by restriction digest. The following abbreviations are used to associate primary accession numbers given in the feature table with their source databases: Em., EMBL; Sw., SWISSPROT; Tr., TrEMBL; Wp., WORMPEP; Information on the WORMPEP database can be found at http://www.sanger.ac.uk/Projects/C_elegans/wormpep This sequence was generated from part of bacterial clone contigs of human chromosome 1, constructed by the Sanger Centre Chromosome 1 Mapping Group. Further information can be found at http://www.sanger.ac.uk/HGP/Chrl Rp11-430G6 is from the library RPCT-11.2 constructed by the group of Pieter de Jong. For further details see http://www.chori.org/bacpac/home.htm VECTOR: pBACes.6 IMPORTANT: This sequence is not the entire insert of clone Rp11-430G6 It may be shorter because we sequence overlapping sections only once, except for a short overlap. The true right end of clone Rp11-430G6 is at 165329 in this sequence. The true right end of clone Rp11-331H2 is at 2000 in this sequence.
BASE COUNT	51144 a 30897 c 31439 g 51849 t
ORIGIN	
FEATURES	
source	
1..165329	
/organism="Homo sapiens"	
/mol_type="genomic DNA"	
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Query Match      100.0%; Score 1000; DB 9; Length 165329;
Best Local Similarity 100.0%; Pred. No. 1.1e-185;
Matches 1000; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 AGTTCAAGACCGCTGAGCAACATGGTGAACCCCATCTCTACTAAAAATACAAAATTA 60
Db 71683 AGTTCAAGACCGCTGAGCAACATGGTGAACCCCATCTCTACTAAAAATACAAAATTA 71742
QY 61 GACAGCGCTGGTGATACAGCCCTGTATCCAGCTACTTCGGAGGCGGAGGAGAAAT 120
Db 71743 GACAGCGCTGGTGATACAGCCCTGTATCCAGCTACTTCGGAGGCGGAGGAGAAAT 71802
QY 121 CACTTGAACCTGCTGGGGTGGAGGTTCGGGGAGCAAGATCATGCGATGCCATCCAGC 180
Db 71803 CACTTGAACCTGCTGGGGTGGAGGTTCGGGGAGCAAGATCATGCGATGCCATCCAGC 71862
QY 181 CCAGGCAACAAGCGGAATGTCTCATCTCAGAAAAAAGGCAATTTATATATATATA 240
Db 71863 CCAGGCAACAAGCGGAATGTCTCATCTCAGAAAAAAGGCAATTTATATATATA 71922
QY 241 TATATATATATACACACACACATATATATATATATATATATATATATATATATA 300
Db 71923 TATATATATATACACACACACATATATATATATATATATATATATATATATATA 71982
QY 301 TATATACACATATATACACATATATATACACATATATGTACACATATATATACACATTA 360
Db 71983 TATATACACATATATACACATATATATATATATATATATGTACACATATATATACACATTA 72042
QY 361 TGTATACACATATATACACATATATACACATATATATACACATATATATATATATA 420
Db 72043 TGTATACACATATATACACATATATATACACATATATATATATATATATATATATA 72102
QY 421 CACATATATACACATATATACACATATATACACATATATATATATATATATATACATA 480
Db 72103 CACATATATACACATATATATACACATATATATATATATATATATATATATATATACATA 72162
QY 481 TATATATATATACACACATATATATACACATATATATACACATATATATATATATATATAT 540
Db 72163 TATATATATATACACACATATATATATACACATATATATATATATATATATATATATATAT 72222
QY 541 ACATATATATACACATATATATACACATATATATATATATATATATATATATATATATATAT 600
Db 72223 ACATATATATACACATATATATATATATATATATATATATATATATATATATATATATATAT 72282
QY 601 ACATATATATATATATATATATATATATATATATATATATATATATATATATATATATATAT 660
Db 72283 ACATATATATATATATATATATATATATATATATATATATATATATATATATATATATATAT 72342
QY 661 ACACATAGATATATATATATATATATATATATATATATATATATATATATATATATATATAT 720
Db 72343 ACACATAGATATATATATATATATATATATATATATATATATATATATATATATATATATAT 72402
QY 721 GCTCAGAGTTTCAAGAGGTAGCAGTTGATTACCACTGGGGATAGAGAAAGAGAGTT 780
Db 72403 GCTCAGAGTTTCAAGAGGTAGCAGTTGATTACCACTGGGGATAGAGAAAGAGAGTT 72462
QY 781 TCACAGCAGTGTATTTGTGAGAGGACATTTTCAGTTGTAGTGGCAATAGTAGGGGAAATAC 840
Db 72463 TCACAGCAGTGTATTTGTGAGAGGACATTTTCAGTTGTAGTGGCAATAGTAGGGGAAATAC 72522
QY 841 ATAAATGTGTATAAAACCTATCTGTAGAGTGTAGTTTAAAGAGGTAAACCTATATATATATATA 900
Db 72523 ATAAATGTGTATAAAACCTATCTGTAGAGTGTAGTTTAAAGAGGTAAACCTATATATATATA 72582
QY 901 TAGTGAAGCAGTGTAAACCTTAAAGGATGGGCCAAGGATTTAAATGTTTATAGGAAGATGG 960
Db 72583 TAGTGAAGCAGTGTAAACCTTAAAGGATGGGCCAAGGATTTAAATGTTTATAGGAAGATGG 72642
QY 961 CTACATGCCAAGCTCAGTGTATGTGGCAGAGGATGGT 1000
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```

RESULT 3  
AC031977/cLOCUS  
DEFINITIONACCESSION  
VERSIONKEYWORDS  
SOURCE

## ORGANISM

REFERENCE  
AUTHORSJOURNAL  
AUTHORSTITLE  
JOURNAL

## COMMENT

AC031977 191699 bp DNA linear HTG 12-APR-2001  
Homo sapiens chromosome 1 clone RP11-288018, WORKING DRAFT  
SEQUENCE, 3 unordered pieces.

AC031977 GI:13194952  
HTG; HTGS\_PHASE1; HTGS\_DRAFT; HTGS\_FULLTOP; HTGS\_ACTIVEFIN.  
Homo sapiens (human)

Homo sapiens  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Primates; Catarrhini; Homnidae; Homo.  
1 (bases 1 to 191699)

Abola, A.P., Bruno, D., Conn, L., Dela Rosa, M., Faulkner, D.,  
Fedorpiel, N., Glukhov, S., Hansen, N., Herman, Z.S., Hyman, R.,  
Mao, J., Marathe, R., Morehouse, A.J., Oefner, P., Palm, C.J.,  
Ramirez, D., Wilhelm, J., Yu, S. and Davis, R.W.

Direct Submission  
Submitted (03-APR-2000) DNA Sequencing and Technology Center,  
Stanford University, 855 California Avenue, Palo Alto, CA 94304,  
USA

On Mar 4, 2001 this sequence version replaced gi:9665085.  
----- Genome Center  
Center: Stanford DNA Sequencing and Technology Development  
Center

Center code: SDSTDC  
Web site: <http://sequence-www.stanford.edu/group/human/>  
Contact: [hum-info@sequence.stanford.edu](mailto:hum-info@sequence.stanford.edu)

----- Project Information  
Center project name: 880  
Center clone name: RP11-288018  
----- Summary Statistics

Sequencing Vector: M13mp18; X02513; 100% of reads  
Chemistry: Dye-primer; 1% of reads  
Assembly program: Phrap; version 0.990319  
Consensus quality: 190680 bases at least Q40  
Consensus quality: 191287 bases at least Q30  
Consensus quality: 191336 bases at least Q20  
Insert size: 19548; agarose-fp  
Quality coverage: 7.9x in Q20 bases; agarose-fp  
Quality coverage: 8.1x in Q20 bases; sum-of-contigs.

\* NOTE: This is a 'working draft' sequence. It currently  
\* consists of 3 contigs. The true order of the pieces  
\* is not known and their order in this sequence record is  
\* arbitrary. Gaps between the contigs are represented as  
\* runs of N, but the exact sizes of the gaps are unknown.  
\* This record will be updated with the finished sequence  
\* as soon as it is available and the accession number will  
\* be preserved.

\* 1 12646: contig of 12646 bp in length  
\* 12647 12746: gap of unknown length  
\* 12747 94961: contig of 82215 bp in length  
\* 94962 95061: gap of unknown length  
\* 95062 191699: contig of 96638 bp in length.

Location/Qualifiers  
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FEATURES  
source







Elhaj, C., Escotto, M., Falls, T., Ferraguto, D., Flagg, N., Ford, J., Foster, P., Frantz, P., Gabisi, A., Gao, J., Garcia, A., Garner, T., Garza, N., Gill, R., Gorrell, J. H., Gujevara, W., Gunaratne, P., Hale, S., Hamilton, K., Harris, C., Harris, K., Hart, M., Havlak, P., Hawes, A., Hernandez, J., Hernandez, O., Hodgson, A., Hogues, M., Holloway, C., Hollins, B., Homs, F., Howard, S., Huber, J., Hulyk, S., Hume, J., Jackson, L. E., Jacobson, B., Jia, Y., Johnson, R., Jolivet, S., Joudah, S., Karlsson, E., Kelly, S., Khan, U., King, L., Korvach, J., Kovar, C., Kratovic, J., Kureshi, A., Landry, N., Leal, B., Lewis, L. C., Lewis, L., Li, J., Li, Z., Lichtarge, O., Lieu, C., Liu, J., Liu, W., Louisghe, H., Lozano, R. J., Lu, X., Lucier, A., Lucier, R., Luna, R., Ma, J., Maheshwari, M., Mapua, P., Martin, R., Martindale, A., Martinez, E., Massey, E., Mawhiney, E., McLeod, M. P., Meador, M., Mei, G., Metzker, M., Miner, G., Miner, Z., Mitchell, T., Mohabbat, K., Morgan, M., Morris, S., Moser, M., Neal, D., Newton, J., Newton, N., Nguyen, A., Nguyen, N., Nguyen, N., Nickerson, E., Nwokenkwo, S., Oguh, M., Okunolu, G., Orangunye, N., Oviedo, R., Pace, A., Payton, B., Peary, J., Perez, L., Peters, L., Pickens, R., Primus, E., Pu, L. L., Quiles, M., Ren, Y., Rives, M., Rojas, A., Rojubenko, I., Roife, M., Ruiz, S., Savery, G., Scherer, S., Scott, G., Shen, H., Shoohtari, N., Sisson, I., Sodergren, B., Sonaike, T., Sparks, A., Stanley, H., Stone, H., Sutton, A., Svatek, A., Tabori, P., Tamerisa, A., Tamerisa, K., Tang, H., Tansey, J., Taylor, C., Taylor, T., Telford, B., Thomas, N., Thomas, S., Usmani, K., Vasquez, L., Vera, V., Villalón, D., Vinson, R., Wall, R., Wang, S., Ward-Moore, S., Warren, R., Washington, C., Washington, S., Williams, G., Williamson, A., Wleczyk, R., Wooden, S., Worley, K., Wu, C., Wu, Y., Wu, Y. F., Zhou, J., Zorrilla, S., Nelson, D., Weinstein, G. and Gibbs, R.

TITLE  
JOURNAL  
REFERENCE  
AUTHORS  
TITLE  
JOURNAL

COMMENT

source

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FEATURES
source
Location/Qualifiers
1. 181946
/organism="Pan troglodytes"
/mol_type="genomic DNA"
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BASE COUNT	49474 a	40705 c	41770 g	48491 t	1506 others
ORIGIN					

Query Match 40.5%; Score 405.2; DB 2; Length 181946;

[illegible]





SOURCE  
ORGANISM  
Homo sapiens (human)  
Eukaryota  
Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.  
1 (bases 1 to 130349)  
Sulston, J.E. and Waterston, R.  
Toward a complete human genome sequence  
Genome Res. 8 (11), 1097-1108 (1998)  
99063792  
MEDLINE  
PUBMED  
9847074  
2 (bases 1 to 130349)  
Pearman, C. and Cotton, M.  
The sequence of Homo sapiens BAC clone RP11-437E9  
Unpublished  
3 (bases 1 to 130349)  
Waterston, R.H.  
Direct Submission  
Submitted (07-OCT-1999) Genome Sequencing Center, Washington  
University School of Medicine, 4444 Forest Park Parkway, St. Louis,  
MO 63108, USA  
4 (bases 1 to 130349)  
Waterston, R.H.  
Direct Submission  
Submitted (23-MAY-2001) Genome Sequencing Center, Washington  
University School of Medicine, 4444 Forest Park Parkway, St. Louis,  
MO 63108, USA  
5 (bases 1 to 130349)  
Waterston, R.H.  
Direct Submission  
Submitted (07-NOV-2001) Department of Genetics, Washington  
University, 4444 Forest Park Avenue, St. Louis, Missouri 63108, USA  
On May 23, 2001 this sequence version replaced gi:13518261.  
-----  
Center: Washington University Genome Sequencing Center  
Center code: WUGSC  
Web site: <http://genome.wustl.edu/gsc>  
Contact: [sapiens@wustl.wustl.edu](mailto:sapiens@wustl.wustl.edu)  
-----  
Summary Statistics  
-----  
Center project name: H\_NH0437E09  
-----

NOTICE: This sequence may not represent the entire insert of this  
clone. It may be shorter because we only sequence overlapping  
clone sections once, or longer because we provide a small overlap  
between neighboring data submissions.

This sequence was finished as follows unless otherwise noted:  
all regions were double stranded, sequenced with an alternate  
chemistry, or covered by high quality data (i.e., phred quality >=  
30); an attempt was made to resolve all sequencing problems, such  
as compressions and repeats; all regions were covered by sequence  
from more than one subclone; and the assembly was confirmed by  
restriction digest.

MAPPING INFORMATION:  
Mapping information for this clone was provided by Dr. John D.  
McPherson, Department of Genetics, Washington University, St. Louis  
MO. For additional information about the map position of this  
sequence, see <http://genome.wustl.edu/gsc>

SOURCE INFORMATION:  
The RPCI-11 human BAC library was made from the blood of one male  
donor, as described by Osoegawa, K., Moon, P.Y., Zhao, B., Prengren, E.,  
Tateno, M., Catanese, J.J. and de Jong, P.J. (1998) An improved  
approach for construction of bacterial artificial chromosome  
libraries. Genomics 51:1-8. The clone may be obtained either from  
Research Genetics, Inc. (<http://www.resgen.com>) or Pieter de Jong  
and coworkers at the Roswell Park Cancer Institute  
(<http://bacpac.med.buffalo.edu>)  
VECTOR: pBAC3.6

NEIGHBORING SEQUENCE INFORMATION:  
The clone sequenced to the left is RP11-289J14, 200 bp overlap; the  
clone sequenced to the right is AC061984. Actual start of this

Clone is at base position 150577 of RP11-289J14; actual end is at  
base position 130349 of RP11-437E9.

FEATURES  
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/db\_xref="taxon:9606"  
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/map="2"  
/clone="RP11-437E9"  
/clone\_lib="RPCI-11"  
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2618..3136  
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3200..3370  
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3522..3856  
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13138..13321  
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13414..13442  
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16183..16266  
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16629..16997  
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17643..17821  
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28282..28808
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28809..29192
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29193..29577
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29628..29895
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32100..32242
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32477..32750
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32848..33164
/rpt_family="Alu"
33165..33302

Query Match 39.5%; Score 395.2; DB 9; Length 130349;
Best Local Similarity 67.8%; Pred. No. 5.7e-68;
Matches 645; Conservative 0; Mismatches 293; Indels 14; Gaps 6;

QY 4 TCAGACAGCAGCTGAGCAACATGGTGAACCCCATCTCTACTATAAATAC--AAAATTAG 61
Db 39934 TCAGACCATCTGGCTACATGGTGAATCCCATCTCTACTATAAATACAAAATTAG 39993
QY 62 ACAGCATGGTGATACACGCTTGTAATCCAGCTACTTCGGAGGCCGAGGAGGAATC 121
Db 39994 CCAGGCGTGGTGGCGGGTGCCTGTAGTCCAGCTACTCCGGAGGCTGAGGAGGAATG 40053
QY 122 ACTTGACCTGCTGGGGTGGAGGTTGGGGGAGCAAGATCATGCTTGCATCCAGCC 181
Db 40054 GGTGTAACC--CGGAGGTGGAGCTTGCATGAGCGAGATGCCACATGCACTCCAGCC 40111
QY 182 CAGGCAACAGAGCGGAATGTCTATCTCAGAAAAAAGGCAATTT--TATATATATA 238
Db 40112 TGGACAAC--GGAGCAAGACTCCATCTATAAATAATATATAAATATATACATAATATA 40170
QY 239 TATATATATATATACACACACACATATATATATATATATATATATATATATATAT 298
Db 40171 TATATAAATATAGACATAAATATATATATATATATATATATATATATATATATAT 40230

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QY 359 TATGTATACACATATATACACATATATACACATATATACACATATATATACACATATA 418
Db 40291 TATATACAAAATATATATAAATATATAAATAATATATAAATATATATACACATATA 40350
QY 419 TACACATATATACACATATATACACATATATACACATATATACACATATATATACATA 478
Db 40351 AATATATATACACATATATATAAATATATAAATAATATATATATATATATAAATATATA 40410
QY 479 CATATATATA--ATATACACATATATATACACATATATACACATATATATACACATA 535
Db 40411 CACATAAATATATGTATGATATATATACATAAATATATATGTATATAATATATACATA 40470
QY 536 TATACACATATATATACACATATATATACACATATATATACATATATATATATATATA 595
Db 40471 AATATATAAGATATATACATAAATATATATAAATATATATATAAATATATATAAATA 40530
QY 596 TATACACATATATATACACATATATATACACATATATATACACATATATATACACATA 655
Db 40531 TATATAAATAGATATATAAATATATATAAATATATATAAATATATATAAATATATAA 40590
QY 656 TATACACACATAGATATATATATATATATATATATATATATATATATATATATATATA 715
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QY 716 TATATGCTCCAGTTCATAAGAGGTAGCAGTTGATTACCACCTGGGGATAGAGGAAGA 775
Db 40651 TAAATATATAAATATATATATAAATATATATAAATATATATAAATATATAAATATA 40710
QY 776 GAGTTTCAGACGAGTGTATTGTGAGAAGGACATTTTCAGGTTGATGGCAATATAGTAGGGA 835
Db 40711 TATATATATAAATATATATAAATATATAAATATATATAAATATATATAAATATAAATA 40770
QY 836 AATACATAAATGTGTAATAAATACCTTATCTGTA---AGGTAGTTAGAGAGTAACTATA 892
Db 40771 TATATATAAATATATAAATATATAAATATATATAAATATATAAATATATAAATATA 40830
QY 893 TATATATATAGTGAAGAGCAGTGTAAACCTTAAAGGATGGGCCCAAGGATTTTAA 944
Db 40831 TATATAAATATATAAATATATATAAATATATAAATATATAAATATATAAATATAAATA 40882

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AC097269 196413 bp DNA linear HTG 20-OCT-2001  
 AC097269/c Pan troglodytes clone RP43-11J15, WORKING DRAFT SEQUENCE, 18  
 LOCUS unordered pieces.  
 DEFINITION Pan troglodytes (chimpanzee)  
 AC097269 1 GI:16117532  
 VERSION HTG; HTGS PHASE1; HTGS DRAFT; HTGS\_FULLTOP.  
 KEYWORDS Pan troglodytes (chimpanzee)  
 SOURCE Pan troglodytes  
 ORGANISM Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Pan.  
 1 (bases 1 to 196413)  
 Muzny, D.M., Adams, C., Adio-Oduola, B., Ali-osman, F.R., Allen, C.,  
 Alsbrooks, S.L., Amaral, H.C., Are, J.R., Banks, T., Barbarella, J.,  
 Benton, J., Bimaga, K., Blankenburg, K., Bonnin, D., Bouck, J.,  
 Bowie, S., Brieva, M., Brown, E., Brown, M., Bryant, N.P., Buhay, C.,  
 Burch, P., Burkett, C., Burrell, K.L., Byrd, N.C., Carron, T.F.,  
 Carter, M., Cavazos, S.R., Chacko, J., Chavez, D., Chen, G., Chen, R.,  
 Chen, Z., Chowdhry, I., Christopoulos, C., Cleveland, C.D., Cox, C.,  
 Coyle, M.D., Dathorne, S.R., David, R., Davila, M.L., Davis, C.,  
 Davy-Carroll, L., Dederich, D.A., Delaney, K.R., Delgado, O.,  
 Denn, A.L., Ding, Y., Dinh, H.H., Douthwaite, K.J., Draper, H.,  
 Dugan-Rocha, S., Durbin, K.J., Earnhart, C., Edgar, D., Edwards, C.C.,  
 Elhaj, C., Escotto, M., Falls, T., Ferraguto, D., Flagg, N., Ford, J.,  
 Foster, P., Frantz, P., Gabisi, A., Gao, J., Garcia, A., Garner, T.,  
 Garza, N., Gill, R., Gorrell, J.H., Guevara, W., Gunaratne, P., Hale, S.,







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repeat_region /rpt_family="AluJ/FRAN" 16532..17094
repeat_region /rpt_family="MLT1F1" 17653..17674
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repeat_region /rpt_family="MER7C" 18650
repeat_region /rpt_family="LIR17" 18999..19103
repeat_region /rpt_family="LIR17" 19283
repeat_region /rpt_family="TAA)n" 20463..20523
repeat_region /rpt_family="A-rich" 21922..22092
repeat_region /rpt_family="MER41B" 2245..22373
repeat_region /rpt_family="MIR" 23956..24058
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repeat_region /rpt_family="MIR" 25149..25213
repeat_region /rpt_family="L2" 25783..26130
repeat_region /rpt_family="MLT1I" 26174..26539
repeat_region /rpt_family="MLT1A1" 28187..28528
repeat_region /rpt_family="AluY" 28792..29263
repeat_region /rpt_family="Charlie8" 30652..30730
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repeat_region /rpt_family="LIME2" 31590..32174
repeat_region /rpt_family="LIME2" 32162..32588
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repeat_region /rpt_family="MIR" 34416..34803
repeat_region /rpt_family="MSTB" 35044..35350
repeat_region /rpt_family="AluSg" 36407..36695
repeat_region /rpt_family="AluSx" 37670..37790
repeat_region /rpt_family="FLAM_A" 38065..38132
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Query Match 39.1%; Score 390.8; DB 9; Length 189662;
Best Local Similarity 65.9%; Pred. No. 4e-67;
Matches 643; Conservative 0; Mismatches 327; Indels 6; Gaps 5;

Qy 1 AGTTCAAGACGAGCTGAGCAACATGGTGAAACCCCATCTCTACTTAAATAT-ACAAAATT 59
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Qy 60 AGACAGGCATGGTGATACACGCGCTGTAATCCAGCTACTTTCGGA-GGCCGAGGCGAGAGA 118
Db 65853 AGCTGGGTGGTGCGAGGTGCTATCTTAATCCAGCTACTCTGGAGGCTGAGCGAGAGA 65912

Qy 119 ATCACTTGAACCTGCTGGGGTGGAGGTTCGGGGAG-CAAGATCATGCCATTGCACCTCC 177
Db 65913 ATTGCTTTGAACC--CAGGAGGTGGAGCTTCAGTGAGCGAGATCGGCCATTGCACCTCC 65970

Qy 178 AGCCGAGCAACAGAGCGGAAATGTCTCTCGAAGAAAAAAGGCAATTTATATATAT 237
Db 65971 AGCCTGGGCAACAGAGCAAACTCTGCTCTCAAAAAAATTTTATAATATATATAT 66030

Qy 238 ATATATATATATATACACACACACATATATATATATATATATATATATATATATAT 297
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Qy 298 ACATATATACATATATATACATATATATATATATATATATATATATATATATATATAT 357
Db 66091 ATATATATATATATATATATATATATATATATATATATATATATATATATATATAT 66150

Qy 358 ATATGTATACATATATATACATATATATATATATATATATATATATATATATATATAT 417
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Qy 418 ATACATATATATATATATATATATATATATATATATATATATATATATATATATATAT 477
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Qy 478 ACATATATATATATATATATATATATATATATATATATATATATATATATATATATAT 537
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Qy 658 TACACACATAGATATATATATATATATATATATATATATATATATATATATATATATAT 717
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Qy 718 TATGCTCCAGGTTTATAAGAGGTAGCAGTTGATTAACCTGGGGATAGAGGAAAGAGA 777
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Qy 778 GTTTCAGCAGCTGTTGTCGAGAGGACATTTTCAGGTTGATGCGCAATAGTAGGGGAAA 837
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Qy 898 ATATAGTGAAGCGAGTGTAACCTTAAAGAGGTGGGCCAAGGATTTAAATGTTTATAGAAGA 957
Db 66690 ATATATATATATATATATATATATATATATATATATATATATATATATATATATATAT 66749

Qy 958 TGGCTAGATGCCAAA 973
Db 66750 AAAGTATGATCTAGA 66765
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RESULT 11  
AC073310/c  
LOCUS AC073310 125685 bp DNA linear PRI 26-APR-2003  
DEFINITION Homo sapiens BAC clone RP11-49G5 from 7, complete sequence.  
ACCESSION AC073310  
VERSION AC073310.7 GI:13176604  
KEYWORDS HTG.  
SOURCE  
ORGANISM Homo sapiens  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.  
1 (bases 1 to 125685)  
Tulston, J.E. and Waterston, R.  
Toward a complete human genome sequence  
Genome Res. 8 (11), 1097-1108 (1998)  
99063792  
MEDLINE  
9847074  
PUBMED  
2 (bases 1 to 125685)  
Haakenson, W., Hannah, C. and Kang, K.  
The sequence of Homo sapiens BAC clone RP11-49G5  
Unpublished (2001)  
REFERENCE  
3 (bases 1 to 125685)  
Waterston, R.H.  
Direct Submission  
Submitted (14-JUN-2000) Genome Sequencing Center, Washington  
University School of Medicine, 4444 Forest Park Parkway, St. Louis,  
MO 63108, USA  
4 (bases 1 to 125685)  
Waterston, R.H.  
Direct Submission  
Submitted (01-MAR-2001) Genome Sequencing Center, Washington  
University School of Medicine, 4444 Forest Park Parkway, St. Louis,  
MO 63108, USA  
5 (bases 1 to 125685)  
Waterston, R.  
Direct Submission  
Submitted (09-MAY-2001) Department of Genetics, Washington  
University, 4444 Forest Park Avenue, St. Louis, Missouri 63108, USA  
6 (bases 1 to 125685)  
Waterston, R.  
Direct Submission  
Submitted (10-MAY-2001) Department of Genetics, Washington  
University, 4444 Forest Park Avenue, St. Louis, Missouri 63108, USA  
7 (bases 1 to 125685)  
Waterston, R.  
Direct Submission  
Submitted (26-APR-2003) Department of Genetics, Washington  
University, 4444 Forest Park Avenue, St. Louis, Missouri 63108, USA  
On Mar 1, 2001 this sequence version replaced gi:11024926.  
----- Genome Center  
Center: Washington University Genome Sequencing Center  
Center code: WUGSC  
Web site: <http://genome.wustl.edu/gsc>  
Contact: [sapiens@watson.wustl.edu](mailto:sapiens@watson.wustl.edu)  
----- Summary Statistics  
Center project name: H\_NH0049G05  
-----

NOTICE: This sequence may not represent the entire insert of this clone. It may be shorter because we only sequence overlapping clone sections once, or longer because we provide a small overlap between neighboring data submissions.

This sequence was finished as follows unless otherwise noted: all regions were double stranded, sequenced with an alternate chemistry, or covered by high quality data (i.e., phred quality >= 30); an attempt was made to resolve all sequencing problems, such as compressions and repeats; all regions were covered by sequence from more than one subclone; and the assembly was confirmed by restriction digest.

#### MAPPING INFORMATION:

The sequence of this clone was established as part of a mapping and sequencing collaboration between the NHGRI Chromosome 7 Mapping Project (Eric D. Green, Director), John D. McPherson in the Department of Genetics (Washington University), and the Washington University Genome Sequencing Center. For additional information about the map position of this sequence, see <http://www.nhgri.nih.gov/DIR/GRB/CHR7>, send [mailto:egreen@nhgri.nih.gov](mailto:mailto:egreen@nhgri.nih.gov), or see <http://genome.wustl.edu/gsc>

#### SOURCE INFORMATION:

The RP11-11 human BAC library was made from the blood of one male donor, as described by Osoegawa, K., Woon, P.Y., Zhao, B., Frengen, E., Tateno, M., Catanese, J.J. and de Jong, P.J. (1998) An improved approach for construction of bacterial artificial chromosome libraries. *Genomics* 51:1-8. The clone may be obtained either from Research Genetics, Inc. (<http://www.resgen.com>) or Pieter de Jong and coworkers at <http://www.chori.org>  
VECTOR: pBACe3.6

#### NEIGHBORING SEQUENCE INFORMATION:

The clone sequenced to the left is RP11-409M7, 200 bp overlap the clone sequenced to the right is RP11-374N8. Actual start of this clone is at base position 14572 of RP11-409M7 actual end is at base position 125685 of RP11-49G5.

FEATURES	Location/Qualifiers
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	3206..5231
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	7142..7353
repeat_region	/rpt_family="MER1_type"



During sequence assembly data is compared from overlapping clones. Where differences are found these are annotated as variations together with a note of the overlapping clone name. Note that the variation annotation may not be found in the sequence submission corresponding to the overlapping clone, as we submit sequences with only a small overlap as described above.

This sequence has been finished according to sequence map criteria as follows. An attempt is made to resolve all sequencing problems, such as compressions and repeats, but not necessarily within known annotated repeat sequence elements. Where the sequence is ambiguous, there is an annotation using the 'unsure' feature key. The following abbreviations are used to associate primary accession numbers given in the feature table with their source databases:

Em, EMBL; Sw, SWISSPROT; Tr, TREMBL; Wp, WORMPEP; Information on the WORMPEP database can be found at

[http://www.sanger.ac.uk/Projects/C\\_elegans/wormpep](http://www.sanger.ac.uk/Projects/C_elegans/wormpep) This sequence was generated from part of bacterial clone contigs of human chromosome 9, constructed by the Sanger Centre Chromosome 9 Mapping Group. Further information can be found at

<http://www.sanger.ac.uk/HGP/Chr9>

RP11-172F4 is from the library RPCI-11.1 constructed by the group of Pieter de Jong. For further details see

<http://www.chori.org/bacpac/home.htm>

VECTOR: pBAC3.6

IMPORTANT: This sequence is not the entire insert of clone RP11-172F4 it may be shorter because we sequence overlapping sections only once, except for a 100 base overlap.

The true left end of clone RP11-172F4 is at 1 in this sequence. The true left end of clone RP11-240L7 is at 139019 in this sequence. The true right end of clone RP11-392G7 is at 43511 in this sequence.

#### FEATURES

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7313..7619  
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repeat\_region  
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9101..9389  
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9399..9701  
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DB 97558 TTAGCTGGGCGTGGTGGCACATGCTGTAGTCTCAGCTACCTGGAGGCTGAGGCGAGG 97617
QY 118 AATCACTTGACCTGCTGGGGTGGAGGTTGGCGGGAG-CAAGATCATGCCATTGCATTC 176
DB 97618 AATCGCTTGAACC--CGGAGGTTGGAGGTTGGAGTGGCGGAGATCGTCCACTGCATC 97675
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DB 97676 CGGCTTGGCGGAC-AGAGCATGATCGTCTCAAAAAAATAATATATATATATATATAT 97734
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QY 332 ATACATATGATACATATATATATATATATATATATATATATATATATATATATATACAC 391
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VERSION AC004787
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SOURCE Homo sapiens (human)
ORGANISM Homo sapiens
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AUTHORS Loftus,B.J., Kim,U.J., Sneddon,V.P., Kalush,F., Brandon,R., Fuhrmann,J., Mason,T., Crosby,M.L., Barnstead,M., Cronin,L., Delatays Mays,A., Cao,X., Xu,R.X., Kang,H.L., Mitchell,S., Eichler,E.E., Harris,P.C., Venter,J.C. and Adams,M.D.
Genome duplications and other features in 12 Mb of DNA sequence from human chromosome 16p and 16q
Genomics 60 (3), 295-308 (1999)
TITLE
JOURNAL
MEDLINE
PUBMED
REFERENCE 2 (bases 1 to 216021)
AUTHORS Adams,M.D., Loftus,B.J., Zhou,L., Crosby,M., Fuhrmann,J., Mason,T.M., Brandon,R., Kim,U.J., Kerlavage,A.R. and Venter,J.C.
Homo sapiens Chromosome 16 BAC clone CIT987SK-A-952F10
Unpublished
TITLE
JOURNAL
REFERENCE 3 (bases 1 to 216021)
AUTHORS Adams,M.D. and Loftus,B.J.
Direct Submission
TITLE
JOURNAL
REFERENCE 4 (bases 1 to 216021)
AUTHORS Adams,M.D. and Loftus,B.J.
Direct Submission
TITLE
JOURNAL
COMMENT Submitted (24-JUL-1998) The Institute for Genomic Research, 9712 Medical Center Dr., Rockville, MD 20850, USA
Address all correspondence to: Mark Adams The Institute for Genomic Research 9712 Medical Center Dr., Rockville, MD 20850, USA e-mail address: humgen@igr.org. The orientation of the sequence is from SP6 end to T7 end. Genes were identified by a combination of five methods including: XGRAIL (available by anonymous ftp from arthur.epm.ornl.gov), Genefinder (Phil Green, University of Washington), GENSCAN (Chris Burge, http://genome.stanford.edu/~chris/GENSCANW.html) searches of the complete sequence against a peptide database, and the Human gene Index database at TIGR (http://www.tigr.org/tdb/hgi/hgi.html). Genes without peptide homology having spliced EST hits are termed 'Unknown gene product'. Genes encoding tRNAs are predicted by

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VERSION AY007685.2 GI:15991796  
KEYWORDS Homo sapiens (human)  
ORGANISM Homo sapiens  
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AUTHORS Leem,S.H., Londono-Vallejo,J.A., Kim,J.H., Bui,H., Tubacher,E., Solomon,G., Park,J.E., Horikawa,I., Kouprina,N., Barrett,J.C. and Laronov,V.  
TITLE The human telomerase gene: complete genomic sequence and analysis of tandem repeat polymorphisms in intronic regions  
JOURNAL Oncogene 21 (5), 769-777 (2002)  
MEDLINE 11850805  
PUBMED 11850805  
REFERENCE 2 (bases 1 to 92564)  
AUTHORS Londono-Vallejo,J.A.  
TITLE Direct Submission  
JOURNAL Submitted (06-SEP-2000) Centre d'Etudes du Polymorphisme Humain, 27 rue Juliette Dodu, Paris 75010, France  
REFERENCE 3 (bases 1 to 92564)  
AUTHORS Londono-Vallejo,J.A.  
TITLE Direct Submission  
JOURNAL Submitted (10-OCT-2001) Centre d'Etudes du Polymorphisme Humain, 27 rue Juliette Dodu, Paris 75010, France  
REMARK Sequence update by submitter  
COMMENT On Oct 10, 2001 this sequence version replaced gi:12642956.  
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GenCore version 5.1.6  
Copyright (c) 1993 - 2003 Compugen Ltd.

OM nucleic - nucleic search, using sw model

Run on: November 7, 2003, 01:54:16 ; Search time 7837.84 Seconds  
(without alignments)  
10444.222 Million cell updates/sec

Title: US-09-939-209A-3\_COPY\_3000\_5000

Perfect score: 2001

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Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 2888711 seqs, 2045481386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seq length: 0

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Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

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3: gb.in.\*

4: gb.om.\*

5: gb.ov.\*

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7: gb.ph.\*

8: gb.pl.\*

9: gb.pr.\*

10: gb.ro.\*

11: gb.sts.\*

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24: em.ph.\*

25: em.pl.\*

26: em.ro.\*

27: em.sts.\*

28: em.un.\*

29: em.vi.\*

30: em.htg\_hum.\*

31: em.htg\_inv.\*

32: em.htg\_other.\*

33: em.htg\_mus.\*

34: em.htg\_pln.\*

35: em.htg\_rod.\*

36: em.htg\_mam.\*

37: em.htg\_vrt.\*

38: em\_sy.\*

39: em\_htgo\_hum.\*

40: em\_htgo\_mus.\*

41: em\_htgo\_other.\*

Pred. No. is the number of results predicted by chance to have a

score greater than or equal to the score of the result being printed,  
and is derived by analysis of the total score distribution.

# SUMMARIES

Result No.	Score	Query Match	Length	DB	ID	Description
1	2001	100.0	20300	6	AX451337	AX451337 Sequence
2	1999.4	99.9	165329	9	AL583850	AL583850 Human DNA
3	1991.4	99.5	131699	2	AC031977	AC031977 Homo sapi
4	231.2	11.6	234560	2	AC115539	AC115539 Rattus no
5	231.2	11.6	275631	2	AC125563	AC125563 Rattus no
6	196.4	9.8	167891	9	AC013439	AC013439 Homo sapi
7	192	9.6	90429	9	HS55110	AL033408 Human DNA
8	192	9.6	110000	2	AC120598	Continuation (3 of
9	192	9.6	117000	9	AC069222	AC069222 Homo sapi
10	192	9.6	140952	2	AC027182	AC027182 Homo sapi
11	192	9.6	166992	2	AC022883	AC022883 Homo sapi
12	192	9.6	204158	9	AL133383	AL133383 Human DNA
13	191.6	9.6	175447	9	AC091607	AC091607 Homo sapi
14	191.6	9.6	176417	2	AC024160	AC024160 Homo sapi
15	191	9.5	134995	9	AL592156	AL592156 Human DNA
16	190.4	9.5	156087	2	AC022036	AC022036 Homo sapi
17	190.4	9.5	172641	9	AC084235	AC084235 Homo sapi
18	190	9.5	74673	2	AC027814	AC027814 Homo sapi
19	186.8	9.3	188208	9	AL441986	AL441986 Human DNA
20	186.8	9.3	193784	2	AL606964	AL606964 Homo sapi
21	186.8	9.3	193787	9	AC017111	AC017111 Homo sapi
22	186	9.3	167303	9	AP001527	AP001527 Homo sapi
23	186	9.3	169513	9	AP002777	AP002777 Homo sapi
24	186	9.3	189637	2	AC009765	AC009765 Homo sapi
25	186	9.3	192104	9	AC006994	AC006994 Homo sapi
26	185.8	9.3	87845	9	AL583860	AL583860 Human DNA
27	185.2	9.3	60532	9	AL590672	AL590672 Human DNA
28	185.2	9.3	129355	9	AC004853	AC004853 Homo sapi
29	185.2	9.3	170543	9	AC021012	AC021012 Homo sapi
30	185.2	9.3	170544	9	CNS05TCC	AL355093 Human chr
31	185.2	9.3	291371	2	AL355584	AL355584 Homo sapi
32	184.6	9.2	200807	9	AC073278	AC073278 Homo sapi
33	184.2	9.2	211791	9	AL162430	AL162430 Human DNA
34	183.2	9.2	97811	9	AC008634	AC008634 Homo sapi
35	183.2	9.2	115915	9	AC105922	AC105922 Homo sapi
36	182.4	9.1	197168	2	AC055765	AC055765 Homo sapi
37	182.4	9.1	149731	9	HS162013	AL035552 Human DNA
38	181.6	9.1	55469	9	AL445490	AL445490 Human DNA
39	181	9.0	121848	9	AC109456	AC109456 Homo sapi
40	181	9.0	206691	9	AC018752	AC018752 Homo sapi
41	180.6	9.0	132718	9	AC107908	AC107908 Homo sapi
42	180.4	9.0	118097	9	AL589826	AL589826 Human DNA
43	180.2	9.0	62871	9	AL356860	AL356860 Human DNA
44	180.2	9.0	138277	2	AC069107	AC069107 Homo sapi
45	180.2	9.0	137127	2	AC069194	AC069194 Homo sapi

## ALIGNMENTS

RESULT 1  
AX451337  
LOCUS AX451337  
DEFINITION Sequence 3 from Patent WO0216653.  
ACCESSION AX451337  
VERSION AX451337.1 GI:21698388  
KEYWORDS  
SOURCE synthetic construct  
ORGANISM synthetic construct  
artificial sequences.  
REFERENCE  
1  
AUTHORS Levitt,P.R., Mirnics,K., Kodavali,V.C. and Nimgaonkar,V.L.  
TITLE Methods and systems for facilitating the diagnosis and treatment of  
schizophrenia  
JOURNAL Patent: WO 0216653-A 3 28-FEB-2002;

AX451337 Sequence 3 from Patent WO0216653.  
DNA linear PAT 03-JUL-2002









Query Match	99.5%;	Score	1991.4;	DB 2;	Length	191699;	
Best Local Similarity	99.7%;	Pred. No.	0;	Mismatches	6;	Indels	0; Gaps 0;
Matches 1995;	Conservative	0;					
QY	1	ATTGTGTAATATAGCAACCTCCCTTCATCATTTAGGTCTTAGTATATACTACTACTCTCTTAG	60				
Db	9938	ATTGTGTAATATAGCAACCTCCCTTCATCATTTAGGTCTTAGTATATACTACTACTCTCTTAG	9879				
QY	61	AGAAGCTGCTCTTCTTCATCTTAAATAAAGTAATAATTCCTTACCTGTTATTTTAAAG	120				
Db	9878	AGAAGCTGCTCTTCTTCATCTTAAATAAAGTAATAATTCCTTACCTGTTATTTTAAAG	9819				
QY	121	TCATCCGCTGTTTCATTTCTGTTAAAGTCTTATCAAAATTTATCATTTATTTTATTTACAGT	180				
Db	9818	TCATCCGCTGTTTCATTTCTGTTAAAGTCTTATCAAAATTTATCATTTATTTTATTTACAGT	9759				
QY	181	CATGTGCCACATPAACAATGTTTCATCTAGGAGTAAAGAACACAAATGATCTCGGCCCATPAA	240				
Db	9758	CATGTGCCACATPAACAATGTTTCATCTAGGAGTAAAGAACACAAATGATCTCGGCCCATPAA	9699				
QY	241	TATTATAGCTCAGAAATTTCTATTAATCTAGTATATCGAGCCATCATAGTGTAAATGC	300				
Db	9698	TATTATAGCTCAGAAATTTCTATTAATCTAGTATATCGAGCCATCATAGTGTAAATGC	9639				
QY	301	AGGACATTTACCTTTCTATGTTTAGATATGTTAGATACACAAATATATTTCAATTTGTGTTA	360				
Db	9638	AGGACATTTACCTTTCTATGTTTAGATATGTTAGATACACAAATATATTTCAATTTGTGTTA	9579				
QY	361	TAATTTCTCAGATATTCAGTACAGTAACTGCTGTACAGGTTTGTACCTTAGAGTAAT	420				
Db	9578	TAATTTCTCAGATATTCAGTACAGTAACTGCTGTACAGGTTTGTACCTTAGAGTAAT	9519				
QY	421	AGGCTATACCATACAGCTAGGTGTAGTGGCTTAACCATCTAGGTTTGTGTAAAGTA	480				
Db	9518	AGGCTATACCATACAGCTAGGTGTAGTGGCTTAACCATCTAGGTTTGTGTAAAGTA	9459				
QY	481	CATTTCTATGATATTCACCAATGATGAATCACTAACTACATCTCAGAAATGTTTC	540				
Db	9458	CATTTCTATGATATTCACCAATGATGAATCACTAACTACATCTCAGAAATGTTTC	9399				
QY	541	ACTGTTGTAGTGAAGTCCCATGACTATATTTTCTATATATCTGATATTTTGTGATCTG	600				
Db	9398	ACTGTTGTAGTGAAGTCCCATGACTATATTTTCTATATATCTGATATTTTGTGATCTG	9339				
QY	601	CCCATGAGATGATGTGTAAGATCAAGATCCAGATGGGTTCTATCCAGTATAGTAC	660				
Db	9338	CCCATGAGATGATGTGTAAGATCAAGATGGGTTCTATCCAGTATAGTAC	9279				
QY	661	CCACTACACTGTTGGATGTCATATGTTTGTGATTAATATCTCAAGATAGACACC	720				
Db	9278	CCACTACACTGTTGGATGTCATATGTTTGTGATTAATATCTCAAGATAGACACC	9219				
QY	721	TTTCTCAGACACATAAAGATGCTCAATATAAAGTTTGTGAACTGAACGTTATTTGCA	780				
Db	9218	TTTCTCAGACACATAAAGATGCTCAATATAAAGTTTGTGAACTGAACGTTATTTGCA	9159				
QY	781	AATGTAACTGATCCGATTTAAAGAGGACGAAACAGAGGCTCTGGCTCAACACCATACT	840				
Db	9158	AATGTAACTGATCCGATTTAAAGAGGACGAAACAGAGGCTCTGGCTCAACACCATACT	9099				
QY	841	TCTAGAGTCATAAAGAGGTAGCAGTTGATTACACTCTGGCGACAGAGGAAAAAGAGCTTG	900				
Db	9098	TCTAGAGTCATAAAGAGGTAGCAGTTGATTACACTCTGGCGACAGAGGAAAAAGAGCTTG	9039				
QY	901	ACCGCAGGCTACTGTGAGACATTTTCAGGTTGATGGCACAGAAACAGGGGAAATACATAAA	960				
Db	9038	ACCGCAGGCTACTGTGAGACATTTTCAGGTTGATGGCACAGAAACAGGGGAAATACATAAA	8979				
QY	961	TGTGTGGGAATATTCAGTGGTCTGGGATCACTACATAGTAGAATATATAGAGAAAGAG	1020				
Db	8978	TGTGTGGGAATATTCAGTGGTCTGGGATCACTACATAGTAGAATATATAGAGAAAGAG	8919				

RESULT 4

AC111539/c

LOCUS

DEFINITION Rattus norvegicus clone CH230-158A12, WORKING DRAFT SEQUENCE, 3

AC111539

234560 bp

DNA

linear

HTG 13-MAY-2003

unordered pieces.  
AC11539  
VERSION AC11539.5 GI:30579258  
KEYWORDS HTG; HTGS PHASE1; HTGS\_DRAFT; HTGS\_FULLTOP.  
SOURCE Rattus norvegicus (Norway rat)  
ORGANISM Rattus norvegicus  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.

REFERENCE  
AUTHORS 1 (bases 1 to 234560)  
Muzny,D,Marie., Metzker,M, Lee., Abramson,S., Adams,C., Alder,J., Allen,C., Allen,H., Albrooks,S., Amin,A., Anguiano,D., Ayalebechi,V., Ayvagi,A., Ayodeji,M., Baca,E., Baden,H., Baldwin,D., Bandaranaike,D., Barber,M., Barnstead,M., Benahmed,F., Biswal,K., Blair,J., Blankenburg,K., Blyth,P., Brown,M., Bryant,N., Bunay,C., Burch,P., Burrell,K., Calderon,E., Cardenas,V., Carter,K., Cavazos,I., Ceasar,H., Center,A., Chacko,J., Chavez,D., Chen,G., Chen,Y., Chen,Z., Chu,J., Cleveland,C., Cockrell,R., Cox,C., Coyle,M., Cree,A., D'Souza,L., Davila,M.L., Davis,C., Davy-Carroll,L., De Anda,C., Dederich,D., Delgado,O., Denson,S., Deramo,C., Ding,Y., Dinh,H., Divya,K., Draper,H., Dugan-Rocha,S., Dunn,A., Durbin,K., Duval,B., Eaves,K., Egan,A., Escotto,M., Eugene,C., Evans,C.A., Falls,T., Fan,G., Fernandez,S., Finley,M., Flagg,N., Forbes,L., Foster,M., Foster,P., Fraser,C.M., Gabisi,A., Ganta,R., Garcia,A., Garner,T., Garza,M., Gbregregoris,E., Geer,K., Gill,R., Grady,M., Guerra,W., Gunaratne,P., Haaland,W., Hamil,C., Hamilton,C., Hamilton,K., Harvey,Y., Havlak,P., Hawes,A., Henderson,N., Hernandez,J., Hernandez,R., Hines,S., Hladun,S.L., Hodgson,A., Hogues,M., Hollins,B., Howells,S., Hulyk,S., Hume,J., Idlebird,D., Jackson,A., Jackson,L., Jacob,L., Jiang,H., Johnson,B., Johnson,R., Jolivet,A., Karpathy,S., Kelly,S., Kelly,S., Khan,Z., King,L., Kovar,C., Kowis,C., Kraft,C.L., Lebow,H., Levan,J., Lewis,L., Li,Z., Liu,J., Liu,J., Liu,W., Liu,Y., London,P., Longacre,S., Lopez,J., Lorensheva,L., Loulseghe,H., Lozado,R.J., Lu,X., Ma,J., Maheshwari,M., Mahindartne,M., Mahmoud,M., Malloy,K., Mangum,A., Mangum,B., Mapua,P., Martin,K., Martin,R., Martinez,E., Mathewney,S., McLeod,M.P., McNeill,T.Z., Meenen,E., Milosavljevic,A., Miner,G., Minja,E., Montemayor,J., Moore,S., Morgan,M., Morris,K., Morris,S., Munidas,M., Murphy,M., Naif,L., Nankervis,C., Neal,D., Newton,N., Nguyen,N., Norris,S., Nwakoelameh,O., Okwunonu,G., Olarunpungoon,A., Pal,S., Parks,K., Pasternak,S., Paul,H., Perez,A., Perez,L., Pfannkoch,C., Plopper,M., Polindexter,A., Popovic,D., Primus,E., Pu,L.-L., Puzo,M., Quiroz,J., Rachlin,E., Reeves,K., Regier,M.A., Reigh,R., Reilly,B., Reilly,M., Ren,Y., Reuter,M., Richards,S., Riggs,F., Rives,C., Rodkey,T., Rojas,A., Rose,M., Rose,R., Ruiz,S.J., Sanders,W., Savery,G., Scherer,S., Scott,G., Shatsman,S., Shen,H., Shetty,J., Shvartsbeyn,A., Sisson,I., Sitter,C.D., Smajs,D., Sneed,A., Sodergren,E., Song,X.-Z., Sorelle,R., Sosa,J., Steinle,M., Strong,R., Sutton,A., Svatek,A., Taber,P., Taylor,C., Taylor,T., Thomas,N., Thomas,S., Tingey,A., Trejos,Z., Usmani,K., Valas,R., Vera,V., Villasana,D., Waldron,L., Walker,B., Wang,J., Wang,Q., Wang,S., Warren,J., Warren,R., Wei,X., White,P., Williams,G., Willson,R., Wleczyk,R., Wooden,H., Worley,K., Wright,D., Wright,R., Wu,J., Yakub,S., Yen,J., Yoon,L., Yoon,V., Yu,F., Zhang,J., Zhou,X., Zhou,S., Zhao,S., Dunn,D., von Niederhausern,A., Weis,R., Smith,D.R., Holt,R.A., Smith,H.O., Weinstock,G. and Gibbs,R.A.

Direct Submission  
Unpublished  
2 (bases 1 to 234560)  
Worley,K.C.  
Direct Submission  
Submitted (19-FEB-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
3 (bases 1 to 234560)  
Rat Genome Sequencing Consortium.  
Direct Submission  
Submitted (13-MAY-2003) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA

COMMENT  
On May 13, 2003 this sequence version replaced gi:25007559. The sequence in this assembly is a combination of BAC based reads and whole genome shotgun sequencing reads assembled using Atlas (<http://www.hgsc.bcm.tmc.edu/projects/rat/>). Each contig described in the feature table below represents a scaffold in the Atlas assembly (a 'contig-scaffold'). Within each contig-scaffold, individual sequence contigs are ordered and oriented, and separated by sized gaps filled with Ns to the estimated size. The sequence may extend beyond the ends of the clone and there may be sequence contigs within a contig-scaffold that consist entirely of whole genome shotgun sequence reads. Both end sequences and whole genome shotgun sequence only contigs will be indicated in the feature table.

Center: Baylor College of Medicine  
Center code: BCM  
Web site: <http://www.hgsc.bcm.tmc.edu/>  
Contact: hgsc-help@bcm.tmc.edu  
Project Information  
Center project name: GMPF  
Center clone name: CH230-158A12  
Summary Statistics  
Assembly program: Atlas 3.0  
Consensus quality: 212222 bases at least Q40  
Consensus quality: 224241 bases at least Q30  
Consensus quality: 226292 bases at least Q20  
Estimated insert size: 231347; sum-of-contigs estimation  
Quality coverage: 7x in Q20 bases; sum-of-contigs estimation

\* NOTE: Estimated insert size may differ from sequence length (see [http://www.hgsc.bcm.tmc.edu/docs/genbank\\_draft\\_data.html](http://www.hgsc.bcm.tmc.edu/docs/genbank_draft_data.html)).  
\* NOTE: This is a 'working draft' sequence. It currently consists of 3 contigs. The true order of the pieces is not known and their order in this sequence record is arbitrary. Gaps between the contigs are represented as runs of N, but the exact sizes of the gaps are unknown.  
\* This record will be updated with the finished sequence as soon as it is available and the accession number will be processed.

1 231343: contig of 231343 bp in length  
231344 231443: gap of unknown length  
231444 232446: contig of 1003 bp in length  
232447 232546: gap of unknown length  
232547 234560: contig of 2014 bp in length.

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/db\_xref="taxon:10116"  
/clone="CH230-158A12"  
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1. .1537  
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misc\_feature  
3731. .7474  
/notes="wgs\_contig"  
misc\_feature  
225356. .228724  
/notes="wgs\_contig"  
misc\_feature  
230080. .231343  
/notes="wgs\_contig"

BASE COUNT 65515 a 49188 c 46995 g 66014 t 6848 others  
ORIGIN

Query Match 11.6%; Score 231.2; DB 2; Length 234560;  
Best Local Similarity 63.7%; Pred. No. 3.1e-43;  
Matches 503; Conservative 0; Mismatches 238; Indels 49; Gaps 8;  
QY 1182 GAGGATCTATAATGGAATCCAGATCTGCTCTCTTAAGTTCAAGCACTTCCATGAC 1241  
|||||  
Db 7000 GAGGACAGAAATTAATAATCTAGTCTGCTCTGTTAGGTTTAAAGCAATGATG-- 6943  
|||||  
QY 1242 ACCATCTGTTTCTTCCACCTGCACAAATGCAATGAACTCTTATGAAACTGCTGTTCT 1301  
|||||  
Db 6942 -----CTGCCTCTTCCACCTGCACAAATGCAATGATCTCTTATGTAATTGCTGTCTA 6889  
|||||

QY 1302 ATCCCTGGCTAAATGTTGCGAAAAAGATTTAACTTTGGGATTAAGCTATTTGGGTT 1361  
 DB |||||  
 QY 6888 TATTGGGCTAAAGCCCTTCCCAAGAAAGAGACAATCTTTGGGAATAAGATTTGTTGGTTC 6829  
 DB |||||  
 QY 1362 TTCTCTCTACTCTCTTGGGAACAA--GGTTTCTTCCCTCGGCTGAATTAAGTGTGGTATTG 1419  
 DB |||||  
 QY 6828 ATCCCTCACTCTTGGGAAGAAAGGCTCTGTGTCCTCCAGCTAGTTAAGCATGTGACAG 6769  
 DB |||||  
 QY 1420 TTCTTCAGGAAATCAGTATGTCATCCTGCTGCTATCAATATGTCAGGTTGGAGTTC 1479  
 DB |||||  
 QY 6768 TTCTTTCTAG-----AGTGATGTCATCCTGCTGCTTGTCAAGTGTGAGGTCACAGTTC 6716  
 DB |||||  
 QY 1480 CTGATTTATGTCATGTGCCCAAAAGCTTGGTGCMAAGAAATGAGACATATTTCCCAAAAG 1539  
 DB |||||  
 QY 6715 TTGGTTTATGTC-----AGAGCCAAACGTTTCCCAAAAG 5680  
 DB |||||  
 QY 1540 TAAGACATCTAGGG--AAGTCCCTGTTTACCTCTGCTGGTATACAGCATCTCCAGGCCCAT 1598  
 DB |||||  
 QY 6679 CCATGGATACCTGGGAAGTCCCTATGCTGATTTCTCTGGCATACAGCACACTCTGGGCTGCT 6620  
 DB |||||  
 QY 1599 ATCTTTGCTTTTGTAGTCTTAAATCAATAA--CTGAACCTCTCATTTGATGCTAGGCCAT 1656  
 DB |||||  
 QY 6619 CTGTTTGTCTGTTATCTCTTAAAACTTACATCATGAACTCTCATGATGTGAGGCCAC 6560  
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 QY 1657 TGTAGTAAACAATAAAGAGAGGGAGGCTTCTGACAACTGAGAGGAAATTTGCTATCTGA 1716  
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 QY 6559 TGTAGGAGACAATAAAGAGAGATAGAGGTTTTTTCACAGCTTAGAGGAAATTTGATCTGA 6500  
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 QY 6499 GGTAGAGCAAAATAACCTTAAGGCTGAGTTGGCTTCATCTCTGCCCTGGAGAGCTGC 6440  
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 DB |||||  
 QY 6439 GATCTCCCTTTAGATCTGATGAATCTAGAGAAATTTCAAGAGCTGCGCCAGTGCCTT 6380  
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 QY 1834 ATTTCCCTGATTTGATACCGTCAATCTTGGAGAAATGTTTTTTTGTCTCCCTGAGC 1893  
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 QY 6379 ATTTCTTGTGTGTAAGATCACAAAATACTTGAGAAAGATTTCTTT--TTCCACAGGAGA 6322  
 DB |||||  
 QY 1894 AAAGGTTGAAAAATTTGAATTTTACCTAGAGACACACATAGTTCCATCTCTGCTGTGT 1953  
 DB |||||  
 QY 6321 GAATCTGGGGATTTCMAATATATCATGTTACACACACAGCTCCGAGCAGCTGCT 6262  
 DB |||||  
 QY 1954 GGCTGAATGT 1963  
 DB |||||  
 QY 6261 GACTTCATGT 6252  
 DB |||||  
 RESULT 5  
 AC125563/c  
 LOCUS Rattus norvegicus clone CH230-9B12, WORKING DRAFT SEQUENCE, 4  
 DEFINITION  
 AC125563  
 VERSION AC125563.4 GI:24817906  
 KEYWORDS HTG; HTGS\_PHASE1; HTGS DRAFT; HTGS\_FULLTOP.  
 SOURCE Rattus norvegicus (Norway rat)  
 ORGANISM Rattus norvegicus  
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae;  
 Rattus.  
 REFERENCE 1 (bases 1 to 275631)  
 AUTHORS Muzny, D. Marie., Metzker, M. Lee., Abramson, S., Adams, C., Alder, J., Allen, C., Allen, H., Alsbrooks, S., Amin, A., Anguiano, D., Anyalebechi, V., Aoyagi, A., Ayodeji, M., Baca, E., Baden, H., Baldwin, D., Bandaranaike, D., Barber, M., Barnstead, M., Benahmed, F., Biewald, K., Blair, J., Blankenburg, K., Blyth, P., Brown, M., Bryant, N., Buhay, C., Burch, P., Burrell, K., Calderon, E., Cardenas, V., Carter, K., Cavazos, I., Caesar, H., Center, A., Chacko, J., Chavez, D., Chen, G., Chen, R., Chen, Y., Chen, Z., Chu, J., Cleveland, C., Cockrell, R., Cox, C., Coyle, M., Cree, A., D'Souza, L., Davila, M. L., Davis, C., Davy-Carroll, L., De Anda, C., Dederich, D.,

Delgado, O., Denson, S., Deramo, C., Ding, Y., Dinh, H., Divya, K., Draper, H., Dugan-Rocha, S., Dunn, A., Durbin, K., Duval, B., Eaves, K., Egan, A., Escotto, M., Eugene, C., Evans, C.A., Falls, T., Fan, G., Fernandez, S., Finley, M., Flagg, N., Forbes, L., Foster, M., Foster, P., Fraser, C.M., Gabisi, A., Ganta, R., Garcia, A., Garner, T., Garza, M., Gebregorjis, E., Geer, K., Gill, R., Grady, M., Guerra, W., Guevara, M., Gunaratne, P., Haaland, W., Hamil, C., Hamilton, C., Hamilton, K., Harvey, Y., Havlak, P., Hawes, A., Henderson, N., Hernandez, J., Hernandez, R., Hines, S., Hladun, S.L., Hodgson, A., Hogue, M., Hollins, B., Howells, S., Huiyk, S., Hume, J., Idler, D., Jackson, A., Jackson, L., Jacob, L., Jiang, B., Johnson, B., Johnson, R., Jolivet, A., Karpach, S., Kelly, S., Kelly, S., Khan, Z., King, L., Kovac, C., Kowis, C., Kraft, C.L., Lebow, H., Levan, J., Lewis, L., Li, Z., Liu, J., Liu, J., Liu, W., Liu, Y., London, P., Longacre, S., Lopez, J., Lorenshuwa, L., Loulsegod, H., Lozano, R.J., Lu, X., Ma, J., Maheshwari, M., Mahindartne, M., Mahmoud, M., Malloy, K., Mangum, A., Mangun, B., Mapua, P., Martin, K., Martin, R., Martinecz, E., Mathew, S., McLeod, M.P., McNeill, T.Z., Meenen, E., Milosavljevic, A., Minet, G., Minja, E., Montemayor, J., Moore, S., Morgan, M., Morris, K., Morris, S., Munidasa, M., Murphy, M., Nair, L., Nankervis, C., Neal, D., Newton, N., Nguyen, N., Norris, S., Nwankwelu, O., Okunnu, G., Olarnpunsagoon, A., Pal, S., Parks, K., Pasternak, S., Paul, H., Perez, A., Perez, L., Pfamkoch, C., Plopper, F., Poindestre, A., Popovic, D., Primus, E., Pu, L.-L., Puazo, M., Quiroz, J., Rachlin, E., Reeves, K., Regier, M.A., Reigh, R., Reilly, B., Reilly, M., Ren, Y., Reuter, M., Richards, S., Riggs, F., Rives, C., Rodkey, T., Rojas, A., Rose, M., Rose, R., Ruiz, S.J., Sanders, W., Savary, G., Scherer, S., Scott, G., Shatsman, S., Shen, H., Shetty, J., Shvartsbeyn, A., Sisson, I., Sitter, C.D., Smaj, D., Sneed, A., Sodergren, E., Song, X.-Z., Sorelle, R., Sosa, J., Steimle, M., Strong, R., Sutton, A., Svatek, A., Taber, P., Taylor, C., Taylor, T., Thomas, N., Thomas, S., Tingey, A., Trejos, Z., Usmani, K., Vallar, R., Varga, V., Villalana, D., Waldron, L., Walker, B., Wang, J., Wang, O., Wang, S., Warren, J., Warren, R., Wei, X., White, F., Williams, G., Willson, R., Wleczyk, R., Wooden, H., Worley, K., Wright, D., Wright, R., Wu, J., Yakub, S., Yen, J., Yoon, L., Yoon, V., Yu, F., Zhang, J., Zhou, J., Zhou, X., Zhao, S., Dunn, D., von Weinstock, G., Weiss, R., Smith, D.R., Holt, R.A., Smith, H.O., Weinstock, G. and Gibbs, R.A.  
 Direct Submission  
 Unpublished  
 2 (bases 1 to 275631)  
 Worley, K.C.  
 Direct Submission  
 Submitted (29-JUN-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
 3 (bases 1 to 275631)  
 Rat Genome Sequencing Consortium.  
 Direct Submission  
 Submitted (09-NOV-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
 On Nov 9, 2002 this sequence version replaced gi:23096552.  
 The sequence in this assembly is a combination of BAC based reads and whole genome shotgun sequencing reads assembled using Atlas (<http://www.hgsc.bcm.tmc.edu/projects/rat/>). Each contig described in the feature table below represents a scaffold in the Atlas assembly (a 'contig-scaffold'). Within each contig-scaffold, individual sequence contigs are ordered and oriented, and separated by sized gaps filled with Ns to the estimated size. The sequence may extend beyond the ends of the clone and there may be sequence contigs within a contig-scaffold that consist entirely of whole genome shotgun sequence reads. Both end sequences and whole genome shotgun sequence only contigs will be indicated in the feature table.

----- Genome Center  
 Center: Baylor College of Medicine  
 Center code: BCM  
 Web site: <http://www.hgsc.bcm.tmc.edu/>  
 Contact: [hgsc-help@bcm.tmc.edu](mailto:hgsc-help@bcm.tmc.edu)  
 ----- Project Information  
 Center project name: GDBK

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Center clone name: CH230-9B12
----- Summary Statistics
Assembly program: Phrap; version 0.990329
Consensus quality: 228152 bases at least Q40
Consensus quality: 230310 bases at least Q30
Consensus quality: 231872 bases at least Q20
Estimated insert size: 235029; sum-of-contigs estimation
Quality coverage: 7x in Q20 bases; sum-of-contigs estimation
-----
* NOTE: Estimated insert size may differ from sequence length
  (see http://www.hgsc.bcm.tmc.edu/docs/Genbank_draft_data.html).
* NOTE: This is a 'working draft' sequence. It currently
  consists of 4 contigs. The true order of the pieces
  is not known and their order in this sequence record is
  arbitrary. Gaps between the contigs are represented as
  runs of N, but the exact sizes of the gaps are unknown.
* This record will be updated with the finished sequence
  as soon as it is available and the accession number will
  be preserved.
*
* 1 192878: contig of 192878 bp in length
* 192879 192878: gap of unknown length
* 192979 270295: contig of 77317 bp in length
* 270296 270395: gap of unknown length
* 270396 272405: contig of 2010 bp in length
* 272406 272505: gap of unknown length
* 272506 275631: contig of 3126 bp in length.
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*     complement(56153..56817)
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*     site:EcoRI
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Best Local Similarity 63.7%; Pred. No. 3.1e-43;
Matches 503; Conservative 0; Mismatches 236; Indels 49; Gaps 8;

QY 1182 GAGGATCTATAATTGAATCCAGATCTGCTCTCTGTAAAGTTCAGACACTTCCATGAC 1241
DB 147760 GAGGAGCAAGATTAAATCTAAGTCTGCTCTGCTGAGGTTTAAAGACATGATG- 147703

QY 1242 ACCATCTGTTCTTCCACCTGCACATGCAAAATGAACTCTTATGAACCTGCTTTCT 1301
DB 147702 -----CTGCTCTTCCACCTGCACATGCAAAATGAACTCTTATGAACCTGCTTTCTA 147649

QY 1302 ATCTGGGCTAAATGTTGAGAAAAGATTTAATCTTTGGATGAAGGCTATTTTGGGTT 1361
DB 147648 TATTGGGCTAAGCCCTTCCAGAAAGAGACAATCTTTGGAAATGAATGTTTGGTTC 147589

QY 1362 TTCTCTACTCTTGGGAACAA--GGTTTCTTCCCTGGGCTAATTAAGTGTGATTTG 1419
DB 147588 ATCTCTCACTCTTGGAAAGAAAGGCTCTGTGTCCCGAGTAGTAAAGCATGTGACAG 147529

QY 1420 TTCTCCAGGAAATCAGTGTGATGCATCCTGCTGTATCAATATGTCAGGGTGTGAGTTC 1479
DB 147528 TTCTTTAG-----AGTATGATCATCCTGCTCTTGTCAAGTGTGAGGTCAGAGTCC 147476

QY 1480 CTGATTTATGTCATGTGCCCAACAAAGCTTGGTGCAAGAAATTTGGACACATTTCCCAAAAG 1539

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Db 147475 TTGGTTTATGTC-----AGAGCCAAACGCTTTTCCCAAAAG 147440
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QY 1599 ATCTTGTCTTTTGTAGTCTTAAATCAATAA--CTGAATCTCTANTGATGTCTAGGCCAT 1656
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QY 1657 TGTAGTAAACAATAAAGAGAGGAGGCTTCTGACAACTGAGAGAAATTTGTCACTGCA 1716
DB 147319 TGTAGAGACATAAAGAGAGTAGAGGTTTTTTCACAGCTTAGAGAAATTTGTAATCTGA 147260
QY 1717 AGTGTGTGAAGCACAGCCTGGGGCTGAGCCTTGGCTTACATCTGCTCCCAAGTGGAGATC 1776
DB 147259 GGTAGAGCAATATAACCTTAAGGCTGAGTGTGGCTTCATCTGCTCCCTGGAGAGGCTGC 147200
QY 1777 AGTGCCCCCATTTAAACATCTGTAGAACTAAAGAA---CGCAACGCTGCGCAATGACTT 1833
DB 147199 GATCTCCCTTTTGAATCTGTAGCACTCAGAGAAATTTCAAAGACTGCCCCAGTGCCTT 147140
QY 1834 ATTTCCTCGCATTTGATACCGTCAATCTCTGAGAAATGTTTCTTTTGTCTTCTCCCTGAGC 1893
DB 147139 ATTTCCTTGTGTAAAGATCACAAAATATCTTGAGAAAGATTTCTTT--TTACACAGGAGA 147082
QY 1894 AAGAGTTGGAATAATTTGAAATTTTACCTAGAGACACACATAGTTTCACATCTCTGCTGTGT 1953
DB 147081 GAAATCTGGCGGATTTCAAATTAATATCAGTGTACCAACACAGCTCCAGACGCTGCGT 147022
QY 1954 GGCTGAATGT 1963
DB 147021 GACTTCATGT 147012

RESULT 6
AC013439/c
LOCUS Homo sapiens BAC clone RP11-270G18 from 2, complete sequence.
DEFINITION AC013439
ACCESSION AC013439
VERSION AC013439.11 GI:13270751
KEYWORDS HTG.
SOURCE Homo sapiens (human)
ORGANISM Homo sapiens
Molecular Type: genomic DNA
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
Sulston, J.E. and Waterston, R.
Toward a complete human genome sequence
Genome Res. 8 (11), 1097-1108 (1998)
99063792
MEDLINE 9847074
PUBMED
REFERENCE 1 (bases 1 to 167891)
AUTHORS Nguyen, C., Cotton, M., Hawkins, M. and Spalding, L.
TITLE The sequence of Homo sapiens BAC clone RP11-270G18
JOURNAL Unpublished
AUTHORS 3 (bases 1 to 167891)
REFERENCE Waterston, R.H.
TITLE Direct Submission
JOURNAL Submitted (11-NOV-1999) Genome Sequencing Center, Washington
University School of Medicine, 4444 Forest Park Parkway, St. Louis,
MO 63108, USA
4 (bases 1 to 167891)
REFERENCE Waterston, R.
AUTHORS Direct Submission
TITLE Submitted (09-AUG-2001) Department of Genetics, Washington
University, 4444 Forest Park Avenue, St. Louis, Missouri 63108, USA
5 (bases 1 to 167891)
REFERENCE Waterston, R.
AUTHORS Direct Submission
TITLE Submitted (07-NOV-2001) Department of Genetics, Washington
University, 4444 Forest Park Avenue, St. Louis, Missouri 63108, USA

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## COMMENT

On Mar 10, 2001 this sequence version replaced gi:12280930.  
 ----- Genome Center  
 Center: Washington University Genome Sequencing Center  
 Center code: WUGSC  
 Web site: <http://genome.wustl.edu/gsc>  
 Contact: [sapiens@wustl.edu](mailto:sapiens@wustl.edu)  
 ----- Summary Statistics  
 Center project name: H\_NH0270G18  
 -----

NOTICE: This sequence may not represent the entire insert of this clone. It may be shorter because we only sequence overlapping clone sections once, or longer because we provide a small overlap between neighboring data submissions.

This sequence was finished as follows unless otherwise noted:  
 all regions were double stranded, sequenced with an alternate chemistry, or covered by high quality data (i.e., phred quality >= 30); an attempt was made to resolve all sequencing problems, such as compressions and repeats; all regions were covered by sequence from more than one subclone; and the assembly was confirmed by restriction digest.

## MAPPING INFORMATION:

Mapping information for this clone was provided by Dr. John D. McPherson, Department of Genetics, Washington University, St. Louis MO. For additional information about the map position of this sequence, see <http://genome.wustl.edu/gsc>

## SOURCE INFORMATION:

The RPL11 human BAC library was made from the blood of one male donor, as described by Osoegawa, K., Moon, P.Y., Zhao, B., Frengen, E., Tateno, M., Catanese, J.J. and de Jong, P.J. (1998) An improved approach for construction of bacterial artificial chromosome libraries. Genomics 51:1-8. The clone may be obtained either from Research Genetics, Inc. (<http://www.resgen.com>) or Pieter de Jong and coworkers at the Roswell Park Cancer Institute (<http://bacpac.med.buffalo.edu>)

## VECTOR: pBACE3.6

## NEIGHBORING SEQUENCE INFORMATION:

The clone sequenced to the left is AC068833; the clone sequenced to the right is RP11-88120, 200 bp overlap. Actual start of this clone is at base position 1 of RP11-270G18; actual end is at base position 167891 of RP11-270G18.

## FEATURES

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Db 79281 ATGACAAATACCTTAAGATGATCTTCTCAGATATGTTCTCTGTTAGTACACAT 79222

## RESULT 8

AC120598.2

WPCOMMENT

Sequence split into 4 fragments LOCUS AC120598 Accession AC120598

Fragment Name	Begin	End
AC120598.0	1	110000
AC120598.1	100001	210000
AC120598.2	200001	310000
AC120598.3	300001	367307

Continuation (3 of 4) of AC120598 from base 200001 (AC120598 Homo sapiens, \*\*\* SEQUENCIN

Query Match 9.6%; Score 192; DB 2; Length 110000;

Best Local Similarity 70.7%; Pred. No. 4.2e-34;

Matches 328; Conservative 0; Mismatches 125; Indels 11; Gaps 5;

QY 117 TAAGTCATCGGTGTTTCATTCTGTTAAAGTCTTATCAATTTATCATTTATTTATTTA 176

Db 94957 TAATCAGCATATATCTCAACATGTTTATTTTACTGMAAAAGTCCAGGTACTATG 95016

QY 177 CAGTCATGTCACATACAAATGTTTCAGTCAGGATAGACACAAATGTA--TCTGCC 234

Db 95017 TAGTCATGAGTGCATATATGATGCTTCAGTCAGAAAGGACCATATTTATCTAGTGTGC 95076

QY 235 CCATAATATTATA---AGCTGAGAAATTTCTATTAACCTAGTGTATATCGCAGCCATCAT 290

Db 95077 CCATGAGATATATGAGGAGCTGAAATTTCTGTACCTAGTGTAGGCTCTAGCCATCAT 95136

QY 291 AG---TGTAATGAGACATATACCTTTCTATGTTAGTAT--GTTAGATACAAATAT 346

Db 95137 ATGACTGTGTGCAATGTCATATCTTTCTATGTTTAGATATATTTAGATACAAATAT 95196

QY 347 ATTTCTATGTTTATATTTCTTACAGTATTCAGTACATACATGCTGTACAGTTTGT 406

Db 95197 TTACCATTTGTTTACAGTTTGCCTACAGCATCTAGTACAGTACATGCTGTATAGTTGT 95256

QY 407 AACCTAGGATATAGCTTACCTACATACAGTGTAGTGTAGTGTAGTGTATACCATCTA 466

Db 95257 AGCTTAGGATATATAGGCTATGCGCTAGTGTAGTGTAGTGTAGTGTAGTGTATACCATCTA 95315

QY 467 GGTTCGTGTAAGTACATTTCTATGATATCCCAATGATGAATCACTACATACATTT 526

Db 95316 GGTTCGTGTAAGTACATCTCATGATGTTGTCAATGACAAATCAGTTAATGGTGCATC 95375

QY 527 TCTCAGATGTTTCACTGTGTGAGTGTACCCATGACCTATATTT 570

Db 95376 TCTCAGACATATTTCTCTGTTGTTAAATGACATATACTATCTT 95419

RESULT 9

AC069222

LOCUS 117000 bp DNA linear PRI 29-MAR-2002

DEFINITION Homo sapiens 3 BAC RP11-38323 (Roewell Park Cancer Institute Human

BAC Library) complete sequence.

ACCESSION AC069222

VERSION AC069222.23 GI:19033388

KEYWORDS HTG.

SOURCE Homo sapiens (human)

ORGANISM Homo sapiens

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

1 (bases 1 to 117000)

Muzny,D.M., Adams,C., Aye,J.R., Ayele,M., Allen,C.,

Alsbrooks,S.L., Amarante,H.C., Are,J.R., Ayale,M., Banks,T.,

Barbata,J., Benton,J., Bimaga,K., Blankenburg,K., Bonnin,D.,

Buck,J., Bowie,S., Brieva,M., Brown,E., Brown,M., Bryant,N.P.,

Bunay,C., Burch,P., Burkett,C., Burrell,K.L., Byrd,N.C.,

Carroll,T.F., Carter,M., Cavazos,S.R., Chacko,J., Chavez,D.,

Chen,G., Chen,R., Chen,Z., Chowdhry,I., Christopoulos,C.,

Cleveland,C.D., Cox,C., Coyle,M.D., Dathorne,S.R., David,R.,

Davila,M.L., Davis,C., Davy-Carroll,L., Dederich,D.A.,

Delaney,K.R., Delgado,O., Denn,A.L., Ding,Y., Dinh,H.H.,  
Douthwaite,K.J., Draper,H., Dugan-Rocha,S., Durbin,K.J.,  
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Moore,S., Morgan,M., Moorish,T., Morris,S., Moser,M., Neal,D.,  
Nelson,D., Newton,J., Newton,N., Nguyen,A., Nguyen,N., Nguyen,N.,  
Nickerson,E., Nwokenko,S., Ogih,M., Okwuonu,G., Oragunye,N.,  
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Taylor,T., Teifrod,B., Thomas,N., Thomas,S., Usmani,K., Vasquez,L.,  
Vera,V., Vallalton,D., Vinson,R., Wang,Q., Wang,S., Ward-Moore,S.,  
Warren,R., Washington,C., Watlington,S., Williams,G.,  
Williamson,A., Wleczek,R., Wooden,S., Worley,K., Wu,C., Wu,Y.,  
Wu,Y.F., Zhou,J., Zorrilla,S., Naylor,S.L., Weinstein,G. and  
Gibbs,R.

## Direct Submission

Unpublished

2 (bases 1 to 117000)

Worley,K.C.

Direct Submission

Submitted (22-MAY-2000) Human Genome Sequencing Center, Department

of Molecular and Human Genetics, Baylor College of Medicine, One

Baylor Plaza, Houston, TX 77030, USA

3 (bases 1 to 117000)

Worley,K.C.

Direct Submission

Submitted (28-FEB-2002) Human Genome Sequencing Center, Department

of Molecular and Human Genetics, Baylor College of Medicine, One

Baylor Plaza, Houston, TX 77030, USA

4 (bases 1 to 117000)

Worley,K.C.

Direct Submission

Submitted (01-MAR-2002) Human Genome Sequencing Center, Department

of Molecular and Human Genetics, Baylor College of Medicine, One

Baylor Plaza, Houston, TX 77030, USA

5 (bases 1 to 117000)

Worley,K.C.

Direct Submission

Submitted (29-MAR-2002) Human Genome Sequencing Center, Department

of Molecular and Human Genetics, Baylor College of Medicine, One

Baylor Plaza, Houston, TX 77030, USA

On Mar 1, 2002 this sequence version replaced gi:18958589.

INFORMATION: <http://www.hgsc.bcm.tmc.edu/> or email

gc-help@bcm.tmc.edu

CLONE LENGTH: This sequence does not necessarily represent the  
entire insert of this clone. Overlapping regions of clones are only  
sequenced and submitted once, so the sequence for the remainder of  
the insert may be found in the record for the adjacent clones.  
Overlapping clones are noted at the beginning and end of the  
Features listing.

## ANNOTATION OF FEATURES:

STSs are identified using ePCR (Genome Res. 7:541-550) searches  
of a local database that includes entries from dbSTS, GDB, and  
local mapping efforts.



AC027182/c  
LOCUS AC027182 140952 bp DNA linear HTG 28-MAR-2000  
DEFINITION Homo sapiens chromosome 1 clone RP11-154119 map 1, WORKING DRAFT  
SEQUENCE, 39 unordered pieces.  
ACCESSION AC027182  
VERSION HTG; HTGS PHASE1; HTGS\_DRAFT.  
KEYWORDS  
SOURCE Homo sapiens (human)  
ORGANISM  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.  
REFERENCE 1 (bases 1 to 140952)  
AUTHORS Birren, B., Linton, L., Nusbaum, C. and Lander, E.  
TITLE Homo sapiens chromosome 1, clone RP11-154119  
JOURNAL Unpublished  
REFERENCE 2 (bases 1 to 140952)  
AUTHORS Birren, B., Linton, L., Nusbaum, C., Lander, E., Abraham, H., Allen, N.,  
Anderson, S., Baldwin, J., Barna, N., Bastien, V., Beda, F.,  
Boguslavsky, L., Bouckgalter, B., Brown, A., Burkett, G.,  
Campotiano, A., Castle, A., Choepel, Y., Colangelo, M., Collins, S.,  
Collamore, A., Cooke, P., DeArrellano, K., Dewar, K., Diaz, J. S.,  
Dodge, S., Domino, M., Doyle, M., Ferreira, P., FitzHugh, W., Gage, D.,  
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McCarthy, M., McEwan, P., McGurk, A., McKernan, K., McPheeters, R.,  
McLirim, J., Meneus, L., Mihova, T., Miranda, C., Mlegha, V., Morrow, J.,  
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Roy, A., Santos, R., Schauer, S., Severy, P., Spencer, B.,  
Stange-Thomann, N., Stojanovic, N., Subramanian, A., Talamas, J.,  
Testaye, S., Theodore, J., Tirrell, A., Travers, M., Trigilio, J.,  
Vassiliev, H., Viel, R., Vo, A., Wilson, B., Wu, X., Wyman, D., Ye, W. J.,  
Young, G., Zainoun, J., Zimmer, A. and Zody, M.  
Direct Submission  
Submitted (28-MAR-2000) Whitehead Institute/MIT Center for Genome  
Research, 320 Charles Street, Cambridge, MA 02141, USA  
All repeats were identified using RepeatMasker:  
Smith, A. F. A. & Green, P. (1996-1997)  
http://ftp.genome.washington.edu/RM/RepeatMasker.html  
----- Genome Center  
Center: Whitehead Institute/ MIT Center for Genome Research  
Center code: WIBR  
Web site: http://www-seq.wi.mit.edu  
Contact: sequence\_submissions@genome.wi.mit.edu  
----- Project Information  
Center project name: L7203  
Center clone name: 154.1.19  
----- Summary Statistics  
Sequencing vector: M13; M77815; 100% of reads  
Chemistry: Dye-terminator Big Dye; 100% of reads  
Assembly program: Phrap; version 0.960731  
Consensus quality: 123230 bases at least Q40  
Consensus quality: 131969 bases at least Q30  
Consensus quality: 134958 bases at least Q20  
Insert size: 157000; agarose-fp  
Insert size: 137152; sum-of-ctgigs  
Quality coverage: 2.6 in Q20 bases; agarose-fp  
Quality coverage: 3.6 in Q20 bases; sum-of-ctgigs  
-----  
\* NOTE: This is a 'working draft' sequence. It currently  
\* consists of 39 contigs. The true order of the pieces  
\* is not known and their order in this sequence record is  
\* arbitrary. Gaps between the contigs are represented as  
\* runs of N, but the exact sizes of the gaps are unknown.  
\* This record will be updated with the finished sequence  
\* as soon as it is available and the accession number will  
\* be preserved.  
\* 1 1256: contig of 1256 bp in length  
\* 1257 1356: gap of 100 bp

1357 2453: contig of 1097 bp in length  
2454 2553: gap of 100 bp  
2554 3853: contig of 1300 bp in length  
3854 3953: gap of 100 bp  
3954 5402: contig of 1449 bp in length  
5403 5503: gap of 100 bp  
5504 7076: contig of 1574 bp in length  
7077 8451: gap of 100 bp  
8452 8551: contig of 1275 bp in length  
8552 9933: gap of 100 bp  
9934 10033: contig of 1382 bp in length  
10034 11519: contig of 1486 bp in length  
11520 14442: contig of 2823 bp in length  
14443 14542: gap of 100 bp  
14543 15852: contig of 1310 bp in length  
15853 15952: gap of 100 bp  
15953 18160: contig of 2208 bp in length  
18161 20873: contig of 2613 bp in length  
20874 23073: gap of 100 bp  
23074 23173: contig of 2100 bp in length  
23174 25043: gap of 100 bp  
25044 25149: gap of 100 bp  
25150 27256: contig of 2107 bp in length  
27257 27356: gap of 100 bp  
27357 29855: contig of 2499 bp in length  
29856 32443: gap of 100 bp  
32444 32543: gap of 100 bp  
32544 36328: contig of 3785 bp in length  
36329 36428: gap of 100 bp  
36429 39039: contig of 2611 bp in length  
39040 42587: gap of 100 bp  
42588 42687: gap of 100 bp  
42688 45620: contig of 2933 bp in length  
45621 45720: gap of 100 bp  
45721 48337: contig of 2617 bp in length  
48338 48437: gap of 100 bp  
48438 51930: contig of 3493 bp in length  
51931 52030: gap of 100 bp  
52031 55883: contig of 3853 bp in length  
55884 55983: gap of 100 bp  
55984 58926: contig of 2943 bp in length  
58927 63146: contig of 4120 bp in length  
63147 63246: gap of 100 bp  
63247 66639: contig of 3393 bp in length  
66640 66733: gap of 100 bp  
66734 70463: contig of 3724 bp in length  
70464 74363: contig of 3800 bp in length  
74364 74463: gap of 100 bp  
74464 79663: contig of 5199 bp in length  
79663 79762: gap of 100 bp  
79763 85047: contig of 5285 bp in length  
85048 85147: gap of 100 bp  
85148 89914: contig of 4767 bp in length  
89915 90014: gap of 100 bp  
90015 94457: contig of 4443 bp in length  
94458 94557: gap of 100 bp  
94558 99499: contig of 4942 bp in length  
99500 104279: contig of 4680 bp in length  
104280 104379: gap of 100 bp  
104380 110964: contig of 6585 bp in length  
110965 111064: gap of 100 bp  
111065 118181: contig of 7117 bp in length  
118182 128265: contig of 9984 bp in length

\* 128266 128365: gap of 100 bp  
 \* 128366 140952: contig of 12587 bp in length.  
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Query Match 9.6%; Score 192; DB 2; Length 140952;  
 Best Local Similarity 73.0%; Pred. No. 4.1e-34;  
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Qy 170 TTATTTACAGTCATGTCACATACCAATGTTTCAGTCAGGATAG--AACACAAATGTA 227  
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 Qy 228 TCTGGCCCCATAATATATATAG--CTGAAATTTCTATTAATACTAGTGATCGCAGC 283  
 Db 10634 TGTGGTCTCTAATAATATATATGGAATCTTAAAGGCTAGTGACATTTAGC 10575  
 Qy 284 CATCATAG-TGTAAATCGAGACATTACCTTTTCTAGTTTAGATGTTAGATACACAA 342  
 Db 10574 CATGGTAACTTCCTTTTCATAGCAATCCCTGTTTCGATGTTTAGATGTTAGGTACACAA 10515  
 Qy 343 ATATATTTTCATGTTGTTATATATTTTCTTACAGTATTCAGTACAGTAACATGCTGTACAGGT 402

Db 10514 GTATTTACCATAGTGTTAACAAGTCTACAGAACTCAGTACTGTAAACATGCTGTACAGGT 10455  
 Qy 403 TTGTAACCTAGAGTAATAGGCTATACCATACAGCTTAGGTAGGTAGTAT----- 457  
 Db 10454 TGGCACCTTACCAAGCAATAGGCCACCATATGCCCTAGGTGTATAGTAGGCACTTAGT 10395  
 Qy 458 -AACCATCTAGGTTTGTGTAAGTACATCTCTATGATTTATCCCAATGATGAATCACCTA 516  
 Db 10394 ACACCATCTAGTTTGTGTAAGGACATTTCTATGATTTAACTGTAATGACAAATCACCTA 10335  
 Qy 517 ACTACACATTTCTCAGAAATGTTTCACTGTTGTGAGTGAGTACCCTACATATATTTTCT 574  
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RESULT 11  
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 ACCESSION AC022883  
 VERSION AC022883.3 GI:7249283  
 KEYWORDS HTG; HTGS PHASE1; HTGS DRAFT.  
 SOURCE Homo sapiens (human)  
 ORGANISM Homo sapiens  
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.  
 1 (bases 1 to 166992)  
 Birren, B., Linton, L., Nusbaum, C. and Lander, E.  
 Homo sapiens, clone RP11-447017  
 Unpublished  
 2 (bases 1 to 166992)  
 Birren, B., Linton, L., Nusbaum, C., Lander, E., Abraham, H., Allen, N.,  
 Anderson, S., Baldwin, J., Barna, N., Beckerly, R., Beda, F.,  
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 Norman, C. H., O'Connor, T., O'Donnell, P., Oliver, T. M., Peterson, K.,  
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 Stojanovic, N., Subramanian, A., Talamas, J., Tesfaye, S., Theodore, J.,  
 Tirrell, A., Vassiliev, H., Viel, R., Vo, A., Wu, X., Wyman, D., Ye, W. J.,  
 Zimmer, A. and Zody, M.  
 Direct Submission

TITLE  
 JOURNAL  
 COMMENT  
 Submitted (06-FEB-2000) Whitehead Institute/MIT Center for Genome  
 Research, 320 Charles Street, Cambridge, MA 02141, USA  
 On Mar 16, 2000 this sequence version replaced gi:6978240.  
 All repeats were identified using RepeatMasker:  
 Smit, A.F.A. & Green, P. (1996-1997)  
 http://ftp.genome.washington.edu/RM/RepeatMasker.html

----- Genome Center  
 Center: Whitehead Institute/ MIT Center for Genome Research  
 Center code: WIBR  
 Web site: http://www-seq.wi.mit.edu  
 Contact: sequence\_submissions@genome.wi.mit.edu  
 ----- Project Information  
 Center project name: L6218  
 Center clone name: 447 O.17  
 ----- Summary Statistics  
 Sequencing vector: M13; M77815; 100% of reads  
 Chemistry: Dye-terminator Big Dye; 100% of reads  
 Assembly program: Phrap; version 0.960731  
 Consensus quality: 152238 bases at least Q40  
 Consensus quality: 159904 bases at least Q30  
 Consensus quality: 162428 bases at least Q20  
 Insert size: 170000; agarose-fp

Insert size: 164492; sum-of-contigs  
 Quality coverage: 3.5 in Q20 bases; agarose-fp  
 Quality coverage: 3.6 in Q20 bases; sum-of-contigs

NOTE: This is a 'working draft' sequence. It currently consists of 26 contigs. The true order of the pieces is not known and their order in this sequence record is arbitrary. Gaps between the contigs are represented as runs of N, but the exact sizes of the gaps are unknown. This record will be updated with the finished sequence, as soon as it is available and the accession number will be preserved.

1 1279: contig of 1279 bp in length  
 \* 1280 1379: gap of 100 bp  
 \* 1380 3074: contig of 1695 bp in length  
 \* 3075 3174: gap of 100 bp  
 \* 3175 4899: contig of 1725 bp in length  
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 \* 5000 7406: contig of 2407 bp in length  
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 \* 7507 9444: contig of 1938 bp in length  
 \* 9445 9544: gap of 100 bp  
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 \* 11703 14743: contig of 2941 bp in length  
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 \* 14844 18243: contig of 3400 bp in length  
 \* 18244 18343: gap of 100 bp  
 \* 18344 19862: contig of 1519 bp in length  
 \* 19863 23231: contig of 3269 bp in length  
 \* 23232 23331: gap of 100 bp  
 \* 23332 26785: contig of 3454 bp in length  
 \* 26786 30112: contig of 3227 bp in length  
 \* 30113 30212: gap of 100 bp  
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 \* 33207 33306: gap of 100 bp  
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 \* 37774 41451: contig of 3678 bp in length  
 \* 41452 46864: contig of 5313 bp in length  
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 \* 52871 52970: gap of 100 bp  
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 \* 61003 61102: gap of 100 bp  
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 \* 88725 88824: gap of 100 bp  
 \* 88825 100892: contig of 12068 bp in length  
 \* 100893 100992: gap of 100 bp  
 \* 100993 114691: contig of 13699 bp in length  
 \* 114692 114791: gap of 100 bp  
 \* 114792 131880: contig of 17089 bp in length  
 \* 131881 131980: gap of 100 bp  
 \* 131981 147366: contig of 15386 bp in length  
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Query Match 9.6%; Score 192; DB 2; Length 166992;  
 Best Local Similarity 70.7%; Pred. No. 4e-34;  
 Matches 328; Conservative 0; Mismatches 125; Indels 11; Gaps 5;  
 QY 117 TAAGTCATCGGTGTTTCATCTCTGTTAAAGTCTTTATCACAATTTATCATTTATTTATTTA 176  
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 QY 177 CAGTCATGTCACATACCAATGTTTCAGTCAGGATAGAACACAATGTA--TCTGGCC 234  
 DB 638 TAGTCATGAGTGCATTAATGCTTCAGTCACCAAGACCATATTTATCTAGTGGTC 697  
 QY 235 CCATATATTATA----AGCTGAGAAATTTCTATTAACTAGTCATATCGCAGCCATCAT 290  
 DB 698 CCATGAGATATAATGAGCTGAAAATTCCTGTACCTAGTGAAGTCTTAGCCATCAT 757  
 QY 291 AG---TGTATGAGGACATACCTTTCTATGTTAGATAT-GTTAGATACACAATAT 346

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Db      758 ATGACTGGTGGCAATGCAATATCTTTCTAAGTTTATAGATATATTAGATACCAATAC 817
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Qy      467 GGTGTTGTGTAAGTACATCTTATGATATATCCACATGATGATGATGATGATGATGATGATGAT 526
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RESULT 12
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LOCUS   Human DNA sequence from clone GSI-204112 on chromosome 1. Contains
DEFINITION   ESTs, STSs and GSSs. Contains a novel gene, complete sequence.
ACCESSION   AL133383
VERSION     AL133383.10 GI:10119658
KEYWORDS    HTG.
SOURCE      Homo sapiens (human)
ORGANISM    Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 204158)
Direct Submission
Submitted (05-JAN-2001) Sanger Centre, Hinxton, Cambridgeshire,
CB10 1SA, UK. E-mail enquiries: humquery@sanger.ac.uk
Requests: clonerquest@sanger.ac.uk
On Sep 12, 2000 this sequence version replaced gi:10086005.
During sequence assembly data is compared from overlapping clones.
Where differences are found these are annotated as variations
together with a note of the overlapping clone name. Note that the
variation annotation may not be found in the sequence submission
corresponding to the overlapping clone, as we submit sequences with
only a small overlap as described above.
The following abbreviations are used to associate primary accession
numbers given in the feature table with their source databases:
Em:, EMBL; Sw:, SWISSPROT; Tr:, TREMBL; Wp:, WORMPEP; Information
on the WORMPEP database can be found at
http://www.sanger.ac.uk/Projects/C_elegans/wormpep/GSI-204112
This sequence is the entire insert of clone GSI-204112 This
sequence has been finished according to sequence map criteria as
follows. An attempt is made to resolve all sequencing problems,
such as compressions and repeats, but not necessarily within known
annotated repeat sequence elements. Where the sequence is
ambiguous, there is an annotation using the 'unsure' feature key.
This sequence was generated by the Sanger Centre from part of a
human chromosome 1 bacterial clone contig constructed by John
Cartpen et al, NHGRI, NIH. Further information can be found at
http://www.sanger.ac.uk/HGP/Chrl.
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## FEATURES

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2720..2870
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5066..5201
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6972..7446
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7453..8545
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9613..9666
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9916..10048
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11160..11284
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11348..11672
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11707..11794
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/note="match: GSS: Em:AQ283704"
19948..20130
/note="THE1-INTERNAL repeat: matches 1398..1580 of
consensus"
20153..20452
/note="THE1C repeat: matches 79..371 of consensus"
20453..21357
/note="LIP repeat: matches 2739..3657 of consensus"
21330..21610
/note="11M4 repeat: matches 4033..4299 of consensus"
21611..21907
/note="AluY repeat: matches 1..298 of consensus"
21908..22023
/note="11M4 repeat: matches 3915..4033 of consensus"
22053..22549

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repeat_region	/note="LTR44 repeat: matches 1. .519 of consensus" 22814. .22841
repeat_region	/note="14 copies 2 mer tt 92% conserved" 22922. .22969
repeat_region	/note="24 copies 2 mer tt 79% conserved" 24230. .24396
misc_feature	/note="match: GSS: Em:AQ605495" 24332. .24673
repeat_region	/note="MER65-internal repeat: matches 2142. .2492 of consensus" 24674. .24740
repeat_region	/note="MER51-internal repeat: matches 1581. .1971 of consensus" 25037. .25536
repeat_region	/note="LTPA5 repeat: matches 5655. .6143 of consensus" 25537. .25600
repeat_region	/note="32 copies 2 mer aa 82% conserved" 25625. .27912
repeat_region	/note="MER31-internal repeat: matches 3177. .4880 of consensus" 28380. .28818
repeat_region	/note="LTR44 repeat: matches 25. .519 of consensus" 28830. .29139
repeat_region	/note="ALuJb repeat: matches 1. .308 of consensus" 29145. .30686
repeat_region	/note="L1M4 repeat: matches 2328. .3854 of consensus" 30748. .31111
repeat_region	/note="THE1B repeat: matches 1. .364 of consensus" 31112. .32639
repeat_region	/note="THE1B-INTERNAL repeat: matches 1. .1580 of consensus" 32640. .33010
repeat_region	/note="THE1B repeat: matches 1. .364 of consensus" 33591. .33698
repeat_region	/note="MIR repeat: matches 33. .141 of consensus" 34045. .34274
repeat_region	/note="MIR repeat: matches 2. .262 of consensus" 35105. .35568
repeat_region	/note="MLTIC repeat: matches 1. .466 of consensus" 36939. .37047
repeat_region	/note="Tigger3(Golem) repeat: matches 1. .108 of consensus" 37039. .37745
repeat_region	/note="Tigger3(Golem) repeat: matches 2277. .3027 of consensus" 38735. .39129
repeat_region	/note="L2 repeat: matches 1162. .1576 of consensus" 41401. .41633
repeat_region	/note="MER58A repeat: matches 1. .224 of consensus" 42037. .42383
misc_feature	/note="LTR37A repeat: matches 76. .414 of consensus" complement(44109. .44432)
misc_feature	/note="match: GSS: Em:B35062" complement(44858. .45350)
misc_feature	/note="match: GSS: Em:AQ528681" complement(45342. .45500)
repeat_region	/note="match: GSS: Em:AQ149065" 45661. .46730
repeat_region	/note="MER11C repeat: matches 1. .1071 of consensus" 47237. .47615
misc_feature	/note="MSTA repeat: matches 1. .426 of consensus" 47669. .48158
repeat_region	/note="match: GSS: Em:AQ428522" 47693. .48244
repeat_region	/note="MER41A repeat: matches 1. .554 of consensus" 48262. .48461
misc_feature	/note="MIR repeat: matches 53. .262 of consensus" 48927. .49106
repeat_region	/note="match: GSS: Em:AQ075808" 49149. .49332
repeat_region	/note="MIR repeat: matches 4. .192 of consensus" 49401. .49551
repeat_region	/note="LTPA5 repeat: matches 5992. .6143 of consensus" 50633. .50779

/note="L2 repeat: matches 2541. .2703 of consensus"  
 51198. .51291  
 /note="MIR repeat: matches 14. .115 of consensus"  
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Query Match	9.6%;	Score 192;	DB 9;	Length 204158;
Best Local Similarity	73.0%;	Pred. No. 3.9e-34;		
Matches 305;	Conservative	0;	Mismatches 100;	Indels 13; Gaps 4;
Qy 170	TTATTTTACAGTCATGTGGCCACATACAAATGTTTTCAGTCAGGCAGGATAG--AAACAAATGTA 227			
Db 11291	TTTAAATACAGTCATGTGCTGTCATACACCAATTCAGTCATGATGGGCTATACATATGAC 11232			
Qy 228	TCGTGGCCCATATATATTAAG---CTGAGAAATTTCTATTAACTAGTGTGATATGCGACG 283			
Db 11231	TGTTGGTCTCATATAATTAATATGGAACATGAATAATCTTTAGGCCCTAGTGACATTTGACG 11172			
Qy 284	CATCATAAAG-TGTAAATGACAGGACATTAACCTTTTCTATGTTTAGATATATGTATAGATACACAA 342			
Db 11171	CATGGTAACCTCTCTTTTCATAGCATTAACCTGTTTCGATGTTTAGATAGTTTAGGTACACAA 11112			
Qy 343	ATATATTTTCAATGTGTGTTATTAATTTTCTACAGTATTTTCAGTACAGTAAACATGCTGTACAGGT 402			
Db 11111	GTATTTTACCATATAGTGTACAACTGCTACAGAACTCAGTACTGTAAACATGCTGTACAGGT 11052			
Qy 403	TTCTAACTTAGGAGTAAATAGGCTATACCATACAGCTTAGGCTGTAGTAGGCTAT----- 457			
Db 11051	TGSCACCTTACCAGCATAGGCCACACCATATGCCCTAGGTGTATAGTAGGCATCTAGT 10992			
Qy 458	-AACCATCTAGGTTTGTGTAAGTACATCTTATGATATTCCTCCACATGATGAATCACTCA 516			
Db 10991	ACACCATCTAGATTTTGTGTAAGGACATCTTATGATGTTAACTGATTAACAAATCACTCA 10932			
Qy 517	ACTACACATTTTCAGAAATGTTTCACTGTTGTGAAGTGAACCATGACCATATATTTTCT 574			
Db 10931	ATGACAGCTTTTCAGAACATATTCCTATGTTAGTGAACCTGATATATATCATATCAT 10874			

AC091607 175447 bp DNA linear PRI 31-JUL-2002  
 Homo sapiens 3 BAC RP11-118D22 (Roswell Park Cancer Institute Human BAC library) complete sequence.  
 AC091607  
 AC091607.11 GI:20986401  
 HTG.  
 Homo sapiens (human)  
 Homo sapiens  
 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.  
 1 (bases 1 to 175447)  
 RESULT 13  
 AC091607/c  
 LOCUS  
 DEFINITION  
 ACCESSION  
 VERSION  
 KEYWORDS  
 SOURCE  
 ORGANISM  
 REFERENCE  
 AUTHORS



Ma, J., Maheshwari, M., Mapua, P., Martin, R., Martindale, A., Martinez, E., Massey, E., Mawhinney, E., McLeod, M.P., Meador, M., Mei, G., Metzger, M., Miner, G., Miner, Z., Mitchell, T., Monabbat, K., Moore, S., Morgan, M., Moorish, T., Morris, S., Moser, M., Neal, D., Nelson, D., Newton, J., Newton, N., Nguyen, A., Nguyen, N., Nguyen, N., Nickerson, E., Nwokenkwo, S., Ogih, M., Okwuonu, G., Oragunye, N., Oviedo, R., Pace, A., Payton, B., Peery, J., Perez, L., Peters, L., Pickens, R., Primus, E., Pu, L.L., Quiles, M., Ren, Y., Rives, M., Rojas, A., Rojebokan, I., Rolfe, M., Ruiz, S., Savery, G., Scherer, S., Scott, G., Shen, H., Shoshkari, N., Sisson, I., Sodergren, E., Sonaike, T., Sparks, A., Stanley, H., Stone, H., Sutton, A., Svatek, A., Tabor, P., Tamerisa, A., Tamerisa, K., Tang, H., Tansey, J., Taylor, C., Taylor, T., Telford, B., Thomas, N., Thomas, S., Usmani, K., Vasquez, L., Vera, V., Villalón, D., Vinson, R., Wang, Q., Wang, S., Ward-Moore, S., Warren, R., Washington, C., Watlington, S., Williams, G., Williamson, A., Wleczyk, R., Wooden, S., Worley, K., Wu, C., Wu, Y., Wu, Y.F., Zhou, J., Zorrilla, S., Naylor, S.L., Weinstein, G. and Gibbs, R.

Direct Submission  
Unpublished  
2 (bases 1 to 175447)  
Worley, K.C.

Direct Submission  
Submitted (09-MAY-2001) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
3 (bases 1 to 175447)  
Worley, K.C.

Direct Submission  
Submitted (17-MAY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
4 (bases 1 to 175447)  
Worley, K.C.

Direct Submission  
Submitted (20-MAY-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
5 (bases 1 to 175447)  
Worley, K.C.

Direct Submission  
Submitted (31-JUL-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
ON MAY 20, 2002 this sequence version replaced gi:20900906.  
INFORMATION: <http://www.hgsc.bcm.tmc.edu/> or email [gc-help@bcm.tmc.edu](mailto:gc-help@bcm.tmc.edu)

CLONE LENGTH: This sequence does not necessarily represent the entire insert of this clone. Overlapping regions of clones are only sequenced and submitted once, so the sequence for the remainder of the insert may be found in the record for the adjacent clones. Overlapping clones are noted at the beginning and end of the Features listing.

#### ANNOTATION OF FEATURES:

STSs are identified using ePCR (Genome Res. 7:541-550) searches of a local database that includes entries from dbSTS, GDB, and local mapping efforts.

Repeats are identified using RepeatMasker (A. Smit and P. Green, unpublished.) for Human and Mouse sequences.

Genes and region of sequence similarity are identified by BLAST (Nuc. Acids Res. 25:3389-3402) similarity (expect < 1e-34) to the EST and cDNA sequences. Genes demonstrate at least two exons flanked by consensus splice sites that maintained sequence continuity across the splice junctions. Sequences that are not identical matches are annotated as similar.

SEQUENCING READ COVERAGE: Sequencing is completed to a minimum standard of double strand coverage with a minimum of 2 clones and 2 reads with no ambiguities or 2 chemistries with a minimum of 2 clones and 3 reads with no ambiguities. If the sequence quality for a region does not meet this standard, it will be indicated in the

annotation as Low Coverage.

QUALITY OF INDIVIDUAL BASES: This sequence meets stringent quality standards - estimated error rate less than 1 per 10,000 bases. Reports of lowest quality individual bases and measures of base quality are listed below. Description of the metrics can be found at URL: <http://gc.bcm.tmc.edu:8088/quality.info/genbank.annotation.html>.

#### QUALSTAT-REPORT

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	/db_xref="taxon:9606"
	/chromosome="3"
	/clone="RP11-118D22"
	1..2004
misc_feature	/note="overlaps bases 28102..30105 of clone AC125607"
	/function="clone overlap"
	8..44
repeat_region	/rpt_family="CT-rich"
	complement(310..624)
repeat_region	/rpt_family="AluY"
	complement(1313..1614)
repeat_region	/rpt_family="AluSg"
	3179..3490
repeat_region	/rpt_family="AluX"
	3491..3541
repeat_region	/rpt_family="(GGAA)n"
	3756..4049
repeat_region	/rpt_family="LIME"
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repeat_region	/rpt_family="LIMEC"
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repeat_region	/rpt_family="L2"
	4629..4652
repeat_region	/rpt_family="AT-rich"
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	complement(6746..6889)
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repeat_region	/rpt_family="(TA)n"
	12971..13031
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	13053..13346
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	14783..14858
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	14881..15067
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	15177..15274
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19832 19951
/rpt family="FLAM_A"
repeat_region      /rpt family="20291"
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/rpt family="MT1102"
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/rpt family="AluSx"
repeat_region      24263..24367
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complement(24408..24629)
/rpt family="MIR"
repeat_region      complement(25224..25366)
/rpt family="L1PA13"
repeat_region      complement(25748..26050)
/rpt family="AluSx"
repeat_region      complement(26969..27012)
/rpt family="MIR"
repeat_region      27780..27805
/rpt family="(T)n"
repeat_region      complement(28026..28071)

Query Match      9.6%; Score 191.6; DB 9; Length 175447;
Best Local Similarity 72.0%; Pred. No. 4.9e-34;
Matches 324; Conservative 0; Mismatches 109; Indels 17; Gaps 5;

QY 130 TTTCATCTGTAAAGTCTTATCAACAATTAATCAATTTATTTTATTTAGTCATGTGCCA 189
DB 6077 TTCAACACATTTTAGGTATACAACAACATTTTTCATCAATAAATTAAGTAGTCATGAGCCA 6018

QY 190 CATAACAATGTTTCAGTC--AGGGATAGAACAACAAATGATCTGGCCCAATAATATTATA 247
DB 6017 CATAATGACATTTTCAGTCAGATGACATCTGCATACATCATGCTGGTCCATAGATTATA 5958

QY 248 A-----GCTGAGAAATTTCTTAATAGTAGTATCGACGCCATCATAGTGAATGCAGG 303
DB 5957 ATGGTCTGAGAAATTTTATAACTAGTAGTACATCATGATATCAATTAATGTTGTAGTACA 5898

QY 304 A-CATTAACCTTTCTATGTTAGATATGTTAGATACACAAATATATTTTCATGTTGTTATA 362
DB 5897 ACATTAACCTTTTCTGTAT-----TTAGATACACAAATACATTAACATGTTGTTATA 5847

QY 363 ATTTCTACATGATTCAGTACAGTAACTGCTGATCAGGTTTGTAACTAGGAGTAATAG 422
DB 5846 ATTGCTTACATATTTAGTACACATATGCTATAAGGTTTGTAGTTAGTACATATA 5787

QY 423 GCTATACATACAGCTTAGTGTGTAGTAGGCTATTAACATCTAGGTTTGTGTAGTACA 482
DB 5786 GCTCTACATATAGCCTAGGTATGGAATAGGCTAT-ACCATCTAGGTTTGTGTAGTACA 5728

QY 483 TTCTATGATATTCACCAATGATGAATCACTAACTACATCACTTCTCAGAAATGTTTAC 542
DB 5727 TTCTATGATATTCACCAATGATGAATTCCTTAAGGAGGCTTCTCAGAAAGTAACCC 5668

QY 543 TGTTGTGAAGTAGCCCATGACTATATTTTC 572
DB 5667 TGTTGTGAAGTAATGCCACCTAGTGTGC 5638

RESULT 14
AC024160
LOCUS      176417 bp DNA linear HTG 29-MAY-2000
DEFINITION Homo sapiens chromosome 3 clone RP11-294L13 map 3p, WORKING DRAFT
SEQUENCE, 19 unordered pieces.
ACCESSION AC024160
VERSION    AC024160.3 GI-8101179
KEYWORDS   HTG; HTGS_PHASE1; HTGS_DRAFT.
SOURCE     Homo sapiens (human)
ORGANISM   Homo sapiens
            Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi;
            Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE  1 (bases 1 to 176417)
AUTHORS   Bao,J., Bao,Q., Bao,W., Bian,X., Cao,T., Chen,C., Chen,J., Ding,H.,
            Dong,W., Fan,H., Feng,X., Guan,Q., Gu,X., Guo,D., He,L., Hu,S.,
            Huang,F., Jin,Y., Kang,N., Li,C., Li,G., Li,J., Li,L.,
            Li,S., Li,T., Liu,Y., Liu,N., Liu,B., Liu,X., Song,S., Sun,M., Sun,W., Sun,Y.,
            Luo,J., Niu,Y., Qi,Q., Qi,X., Song,S., Sun,M., Sun,W., Sun,Y.,
            Tao,R., Wang,H., Wang,J., Wang,L., Wang,L., Wang,L., Wang,R.,
            Wang,X., Wang,X., Wang,Y., Wu,D., Wu,Q., Xie,P., Xuan,Z., Xue,Y.,
            Yan,C., Yang,X., Yu,B., Zeng,Y., Zhang,G., Zhang,H., Zhang,H.,
            Zhang,L., Zhang,M., Zhang,X., Zhang,X., Zhang,Y., Zhang,Y.,
            Zhang,Z., Zhu,B., Yu,J. and Yang,H.
            Unpublished
            Chromosome 3p genomic sequence
            2 (bases 1 to 176417)
            Chen,C., Hu,S., Dong,W., Zhang,X., Wang,J., Zhang,Y., Zhang,H.,
            Liu,B., Bao,W., Sun,Y., Wu,Q., Wang,H., Yang,X., Cheng,C., Wang,Y.,
            Niu,Y., Qi,X., Li,T., Zhang,H., Liu,N., Wu,D., Yu,B., Fan,H.,
            Liu,Y., Li,G., Li,C., Bao,Q., Bao,J., Wang,X., Song,L., Zhang,L.,
            Guo,D., Huang,F., Zhang,G., Li,J., Bian,X., Zhang,M., Li,L.,
            Feng,X., Yu,J. and Yang,H.
            Direct Submission
            Submitted (25-FEB-2000) Human Genomic Center, Institute of
            Genetics, Chinese Academy of Sciences, Datun Road, Beijing, Beijing
            100101, P.R.China
            On May 29, 2000 this sequence version replaced gi:7644436.
            -----Genome Center
            Center:Beijing Center
            Center code:Beijing
            Website:http://hgci.igtp.ac.cn
            http://www.genomics.org.cn
            Contact:hgci@igtp.ac.cn
            -----Project Information
            Center project name:1# project
            Center clone name: RP11-294L13
            -----Summary Statistics
            Sequencing vector: pUC18; 100% of reads
            Chemistry: Dye-terminator; ET 55% of reads
            Chemistry: Dye-terminator Big Dye; 45% of reads
            Assembly program: Phrap; version 0.990329
            Consensus quality: 165854 bases at least Q40
            Consensus quality: 170551 bases at least Q30
            Consensus quality: 173535 bases at least Q20
            Insert size: 16736; sum-of-contigs
            Quality coverage: 4.26x in Q20 bases;sum-of-contigs
            -----
            * NOTE: This is a 'working draft' sequence. It currently
            * consists of 19 contigs. The true order of the pieces
            * is not known and their order in this sequence record is
            * arbitrary. Gaps between the contigs are represented as
            * runs of N, but the exact sizes of the gaps are unknown.
            * This record will be updated with the finished sequence
            * as soon as it is available and the accession number will
            * be preserved.
            * 1 1505: contig of 1505 bp in length
            * 1506 1605: gap of unknown length
            * 1606 4034: contig of 2429 bp in length
            * 4035 4134: gap of unknown length
            * 4135 5856: contig of 1722 bp in length
            * 5857 5957: gap of unknown length
            * 5957 8671: contig of 2715 bp in length
            * 8671 8771: gap of unknown length
            * 8771 14318: contig of 5547 bp in length
            * 14318 14418: gap of unknown length
            * 14419 18764: contig of 4346 bp in length
            * 18765 18864: gap of unknown length
            * 18865 23344: contig of 4480 bp in length
            * 23345 23445: gap of unknown length
            * 23445 28654: contig of 5210 bp in length
            * 28655 28754: gap of unknown length
            * 28755 34797: contig of 6043 bp in length
            * 34798 34898: gap of unknown length
            * 34899 40772: contig of 5875 bp in length
            * 40773 40873: gap of unknown length
            * 40873 47169: contig of 6296 bp in length
            * 47169 53552: contig of 6284 bp in length

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\* 53553 53652: gap of unknown length  
 \* 53653 contig of 12202 bp in length  
 \* 53654 65954: gap of unknown length  
 \* 53655 80747: contig of 14793 bp in length  
 \* 53656 80748: gap of unknown length  
 \* 53657 92030: contig of 11183 bp in length  
 \* 53658 92031 92130: gap of unknown length  
 \* 53659 109264: contig of 17134 bp in length  
 \* 53660 109364: gap of unknown length  
 \* 53661 123322: contig of 13958 bp in length  
 \* 53662 123422: gap of unknown length  
 \* 53663 146425: contig of 23003 bp in length  
 \* 53664 146426 146525: gap of unknown length  
 \* 53665 176417: contig of 29892 bp in length.

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 /clone="RP11-294L13"

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## misc\_feature

1606. 4034  
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## misc\_feature

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## misc\_feature

40873. 47168  
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## misc\_feature

47269. 53552  
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## misc\_feature

53653. 65854  
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## misc\_feature

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## misc\_feature

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## misc\_feature

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## misc\_feature

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## ORIGIN

Query Match 9.6%; Score 191.6; DB 2; Length 176417;  
 Best Local Similarity 72.0%; Pred. No. 4.9e-34;  
 Matches 324; Conservative 0; Mismatches 109; Indels 17; Gaps 5;

QY 130 TTTCATCTGTTAAGTCTTATACAAATTATCATTTTATTACAGTCAGTCGCA 189  
 Db 150340 TTCAACACTTTAGGTATACAAACAACTTTTCATCATTAATTAAGTAGTCAGGCCA 150399

QY 190 CATAACATGTTTCAGTC--AGGATAGAACACAAATGTATCTGGCCCAATAATTATA 247  
 Db 150400 CATAATGACATTTTCAGTCGATGACATCTGATCATCATGTTGGTCCCAAGATTATA 150459  
 QY 248 A-----GCTGAGAAATTTCTATTAACTAGTATATCGGACCATCATAGTGTAAATGCGAGG 303  
 Db 150460 ATGGTGTCTGAAAAATTTTATAAACTAGTCATCATAGTTATCATATAATGTTGTAGTACA 150519  
 QY 304 A-CATTACCTTTCTATGCTTTTAGATATGTTAGATACACAAATATATTTTCATTGTCTTATA 362  
 Db 150520 ACCATTACCTTTCTGTA-----TTAGATACACAAATATCTTACATTGTGTTATA 150570  
 QY 363 ATTTCTACAGTATTCAGTACAGTAACATGCTGACAGGTTTGTAACTAGGAGTAATAG 422  
 Db 150571 ATTGCTACATAATTTAGTACAGCAATATGCTATAAAGGTTTGTAGCTTAGTACCAATAA 150630  
 QY 423 GCTATACCATACAGCTTAGGTGCTAGTAGCTATACCATCTAGGTTTGTAGTACATA 482  
 Db 150631 GCTTACCATATAGCTTAGGTATGGAATAGGCTAT-ACCATCTAGGTTTGTAGTACATA 150689  
 QY 483 TTCTATGATATTTCCCAATGATGAATCACTAACTACATTCATCTCAGAAATGTTTCCAC 542  
 Db 150690 TTCTATGATATTTCCCAAGATGATGAATGCTTAGGAGGATTTCTCAGAAAGTAACCC 150749  
 QY 543 TGTTGTGAAGTGACCCATGACTATATTTTC 572  
 Db 150750 TGTTGTGAAGATGCACCACTGTAGTTGC 150779

## RESULT 15

## AL592156/c

## LOCUS

## DEFINITION

## Human DNA sequence from clone RP11-49208 on chromosome X, complete

## sequence.

## ACCESSION

## AL592156

## VERSION

## AL592156.4

## KEYWORDS

## HTG.

## SOURCE

## Homo sapiens

## ORGANISM

## Homo sapiens

## REFERENCE

## Direct Submission

## AUTHORS

## TITLE

## JOURNAL

## COMMENT

## On Jul 25, 2001 this sequence version replaced gi:14625708.

## During sequence assembly data is compared from overlapping clones.

## Where differences are found these are annotated as variations

## together with a note of the overlapping clone name. Note that the

## variation annotation may not be found in the sequence submission

## corresponding to the overlapping clone, as we submit sequences with

## only a small overlap as described above.

## This sequence was finished as follows unless otherwise noted: all

## regions were either double-stranded or sequenced with an alternate

## chemistry or covered by high quality data (i.e., phred quality &gt;=

## 30); an attempt was made to resolve all sequencing problems, such

## as compressions and repeats; all regions were covered by at least

## one plasmid subclone or more than one M13 subclone; and the

## assembly was confirmed by restriction digest. The following

## abbreviations are used to associate primary accession numbers given

## in the feature table with their source databases: Em:, EMBL; Sw:,

## SWISSPROT; Tr:, TREMBL; Wp:, WORMPEP; Information on the WORMPEP

## database can be found at

## http://www.sanger.ac.uk/Projects/C\_elegans/wormpep This sequence

## was generated from part of bacterial clone contigs of human

## chromosome X, constructed by the Sanger Centre Chromosome X Mapping

## Group. Further information can be found at

## http://www.sanger.ac.uk/HGP/ChrX

## RP11-49208 is from the library RPCI-11.2 constructed by the group

## of Pieter de Jong. For further details see

## http://www.chori.org/bacpac/home.htm

## VECTOR: pBAC3.6

IMPORTANT: This sequence is not the entire insert of clone RP11-49208. It may be shorter because we sequence overlapping sections only once, except for a 100 base overlap. The true right end of clone RP11-49208 is at 134995 in this sequence. The true right end of clone RP13-46M24 is at 2000 in this sequence.

## FEATURES

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Db 109919 TATGTGACCGTCATATGCCACATACCGGTGTTTGGTCAATGGTGGACCATATATGG 109860

Qy 229 C--TGGCCCCATATATATA---AGCTGAGAAATTTCTATTAACTAGTGATATCGCA- 281
Db 109859 CAGTGGTTGGTAAATCAATATGAGAGCTGAAATTTCTATTGCCCTAGTGTGG 109800

Qy 282 --GCCATCATAGTGTAAATGCGAGACATACCTTTTCTATGTTTATAGATATG-TTAGATAC 338
Db 109799 TGGCCCTCATACATTTAGTGTCAACATTAACCTTTCTATATATAGATATGTTTAGATAC 109740

Qy 339 ACAATATATTTTCATTCGTTATTAATTTCTACAGTATTCAGTACAGTACATGCTGTAC 398
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Qy 399 AGGTTTGTAACTAGGAGTAAATAGGCTATACCATACAGCTTAGGTGTGTAGTGGCTATA 458
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Qy 459 ACCATCTAGGTTTGTGTAGTACATTTCTATGATAT--TCCGCAATGATGAATCACCTA 516
Db 109620 ACCATCTAGGTCGTGTAGTATGCTCTATGATATGCTGTCATACCAACAAATCACCTA 109561

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GenCore version 5.1.6  
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Scoring table: IDENTITY\_NUC  
Gapop 10.0 , Gapext 1.0  
Searched: 2888711 seqs, 20454813386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seq length: 0  
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Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

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- 1: gb\_ba.\*
- 2: gb\_htg.\*
- 3: gb\_in.\*
- 4: gb\_om.\*
- 5: gb\_ov.\*
- 6: gb\_pat.\*
- 7: gb\_ph.\*
- 8: gb\_pl.\*
- 9: gb\_pr.\*
- 10: gb\_ro.\*
- 11: gb\_sts.\*
- 12: gb\_sy.\*
- 13: gb\_un.\*
- 14: gb\_vi.\*
- 15: em\_ba.\*
- 16: em\_fun.\*
- 17: em\_in.\*
- 18: em\_hum.\*
- 19: em\_mu.\*
- 20: em\_om.\*
- 21: em\_or.\*
- 22: em\_ov.\*
- 23: em\_pat.\*
- 24: em\_ph.\*
- 25: em\_pl.\*
- 26: em\_ro.\*
- 27: em\_sts.\*
- 28: em\_un.\*
- 29: em\_vi.\*
- 30: em\_htg\_hum.\*
- 31: em\_htg\_inv.\*
- 32: em\_htg\_other.\*
- 33: em\_htg\_mus.\*
- 34: em\_htg\_pln.\*
- 35: em\_htg\_rtd.\*
- 36: em\_htg\_mam.\*
- 37: em\_htg\_vrt.\*
- 38: em\_sy.\*
- 39: em\_htgo\_hum.\*
- 40: em\_htgo\_mus.\*
- 41: em\_htgo\_other.\*

Pred. No. is the number of results predicted by chance to have a

score greater than or equal to the score of the result being printed,  
and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match %	Length	DB	ID	Description
1	501	100.0	20300	6	AX451337	AX451337 Sequence
2	501	100.0	165329	9	AL583850	AL583850 Human DNA
3	501	100.0	191699	2	AC031977	AC031977 Homo sapi
4	167.8	33.5	207387	2	AC115766	AC115766 Mus muscu
5	138.2	27.6	275631	2	AC125563	AC125563 Rattus no
6	69.4	13.9	2599	9	AK093959	AK093959 Homo sapi
7	69	13.8	3769	9	AK096204	AK096204 Homo sapi
8	59	11.8	234560	2	AC111539	AC111539 Rattus no
9	48	9.6	174328	9	AC100845	AC100845 Homo sapi
10	47.6	9.5	7218	6	I66494	I66494 Sequence 14
11	45.6	9.1	307979	2	AC095234	AC095234 Rattus no
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13	45.4	9.1	134971	2	AC116367	AC116367 Oryza sat
14	45.2	9.0	198464	2	AC022764	AC022764 Homo sapi
15	45.2	9.0	203581	10	AL451076	AL451076 Mouse DNA
16	45.2	9.0	252721	2	AC095946	AC095946 Rattus no
17	44.8	8.9	1122	3	AF020286	AF020286 Dictyoste
18	44	8.8	2753	9	BC051869	BC051869 Homo sapi
19	43.6	8.7	144177	9	AL513487	AL513487 Human DNA
20	43.4	8.7	119171	2	EX247904	EX247904 Danio rer
21	43.4	8.7	124895	8	AC135288	AC135288 Solanum d
22	43.2	8.6	179212	9	AC084877	AC084877 Homo sapi
23	43	8.6	59469	2	AC110024	AC110024 Homo sapi
24	43	8.6	173817	9	AC103834	AC103834 Homo sapi
25	42.8	8.5	125020	9	AF429315	AF429315 Homo sapi
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30	42.2	8.4	59584	2	AC090243	AC090243 Homo sapi
31	42.2	8.4	166315	9	AC078940	AC078940 Homo sapi
32	42	8.4	887	9	HS0801336	AL669860 Mouse DNA
33	42	8.4	83912	8	AP002043	AP002043 Arabidops
34	42	8.4	131275	10	AL669860	AL669860 Mouse DNA
35	42	8.4	150681	9	AP001251	AP001251 Homo sapi
36	42	8.4	192187	3	AC116920	AC116920 Dictyoste
37	42	8.4	201299	5	AL773601	AL773601 Zebrafish
38	42	8.4	250029	3	AE014839	AE014839 Plasmodiu
39	42	8.4	340000	9	AP001679	AP001679 Homo sapi
40	41.6	8.3	146570	3	AC117072	AC117072 Dictyoste
41	41.6	8.3	150351	2	AC102755	AC102755 Mus muscu
42	41.6	8.3	250029	3	AE014820	AE014820 Plasmodiu
43	41.4	8.3	38692	3	AC116919	AC116919 Dictyoste
44	41.4	8.3	100549	9	AL138776	AL138776 Human DNA
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ALIGNMENTS

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LOCUS AX451337  
DEFINITION Sequence 3 from Patent WO0216653.  
ACCESSION AX451337  
VERSION AX451337.1 GI:21698388  
KEYWORDS  
SOURCE synthetic construct  
ORGANISM synthetic construct  
REFERENCE  
AUTHORS  
TITLE  
JOURNAL

AX451337 AX451337 20300 bp DNA linear PAT 03-JUL-2002

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FEATURES
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  Best Local Similarity 100.0%; Pred. No. 1.1e-113;
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AL583850 165329 bp DNA linear PRI 15-NOV-2001
LOCUS Human DNA sequence from clone RP11-430G6 on chromosome 1, complete
DEFINITION sequence.
ACCESSION AL583850
VERSION AL583850.5 GI:16973044
KEYWORDS HTG.
SOURCE Homo sapiens (human)
ORGANISM Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 165329)
Tracey,A.
Direct Submission
Submitted (15-NOV-2001) Wellcome Trust Sanger Institute, Hinxton,
Cambridgeshire, CB10 1SA, UK. E-mail enquiries:
humquery@sanger.ac.uk
On Nov 16, 2001 this sequence version replaced g1:15020514.
During sequence assembly data is compared from overlapping clones.
REFERENCE
AUTHORS
TITLE
JOURNAL
COMMENT

```

Where differences are found these are annotated as variations together with a note of the overlapping clone name. Note that the variation annotation may not be found in the sequence submission corresponding to the overlapping clone, as we submit sequences with only a small overlap as described above.

This sequence was finished as follows unless otherwise noted: all regions were either double-stranded or sequenced with an alternate chemistry or covered by high quality data (i.e., phred quality >= 30); an attempt was made to resolve all sequencing problems, such as compressions and repeats; all regions were covered by at least one plasmid subclone or more than one M13 subclone; and the assembly was confirmed by restriction digest. The following abbreviations are used to associate primary accession numbers given in the feature table with their source databases: Em: EMBL; Sw: SWISSPROT; Tr: TREMBL; Wp: WORMPEP; Information on the WORMPEP database can be found at

[http://www.sanger.ac.uk/Projects/C\\_elegans/wormpep](http://www.sanger.ac.uk/Projects/C_elegans/wormpep) This sequence was generated from part of bacterial clone configs of human chromosome 1, constructed by the Sanger Centre Chromosome 1 Mapping Group. Further information can be found at

<http://www.sanger.ac.uk/HGP/Chr1>

RP11-430G6 is from the library RPCI-11.2 constructed by the group of Pieter de Jong. For further details see

<http://www.chori.org/bacpac/home.htm>

VECTOR: pBAC3.6

IMPORTANT: This sequence is not the entire insert of clone RP11-430G6 it may be shorter because we sequence overlapping sections only once, except for a short overlap.

The true right end of clone RP11-430G6 is at 165329 in this sequence. The true right end of clone RP11-331H2 is at 2000 in this sequence.

#### FEATURES

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1

REFERENCE  
AUTHORS  
TITLE  
JOURNAL  
REFERENCE  
AUTHORS

1. (bases 1 to 207387)  
Birren,B., Nusbaum,C. and Lander,E.  
Mus musculus, clone RP23-82124  
Unpublished  
2. (bases 1 to 207387)  
Birren,B., Linton,L., Nusbaum,C., Lander,E., Ali,A., Allen,N.,  
Anderson,S., Barna,N., Bastien,V., Bloom,T., Boguslavsky,L.,  
Boukhalter,B., Brown,A., Camarata,J., Campopiano,A., Chang,J.,  
Chazaro,B., Choepel,Y., Colangelo,M., Collins,S., Collymore,A.,  
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Hagos,B., Horton,L., Hulme,W., Iliiev,I., Johnson,R., Jones,C.,  
Kamat,A., Karatas,A., Kellis,C., LaRocque,K., Lamazares,R.,  
Landers,T., Lehoczy,J., Levine,R., Lindblad-Toh,K., Liu,G.,  
MacLean,C., Macdonald,P., Major,J., Marquis,N., Matthews,C.,  
McCarthy,M., McEwan,P., McKernan,K., Meldrim,J., Meneus,L.,  
Mihova,T., Mlenga,V., Murphy,T., Naylor,J., Nguyen,C., Nicol,R.,  
Norbu,C., Norman,C.H., O'Connor,T., O'Donnell,P., O'Neill,D.,  
Oliver,J., Peterson,K., Phunkhang,P., Pierre,N., Pollara,V.,  
Raymond,C., Retta,R., Rieback,M., Riley,R., Rise,C., Rogov,P.,  
Roman,J., Rosetti,M., Roy,A., Santos,R., Schauer,S., Schupback,R.,  
Seaman,S., Severy,P., Spencer,B., Stange-Thomann,N., Stojanovic,N.,  
Strauss,N., Subramanian,A., Talamas,J., Tesfaye,S., Theodore,J.,  
Topham,K., Travers,M., Travis,N., Triggilio,J., Vassiliev,H.,  
Viel,R., Vo,A., Wilson,B., Wu,X., Wyman,D., Ye,W.J., Young,G.,  
Zainoun,J., Zembek,L., Zimmer,A. and Zody,M.  
Direct Submission  
Submitted (22-MAR-2003) Whitehead Institute/MIT Center for Genome  
Research, 320 Charles Street, Cambridge, MA 02141, USA  
3. (bases 1 to 207387)  
Birren,B., Nusbaum,C., Lander,E., Abouelleil,A., Allen,N.,  
Anderson,S., Arachchi,H.M., Barna,N., Bastien,V., Bloom,T.,  
Boguslavsky,L., Boukhalter,B., Camarata,J., Chang,J., Choepel,Y.,  
Collymore,A., Cook,A., Cooke,P., Corum,B., Dearellano,K.,  
Diaz,J.S., Dodge,S., Dooley,K., Dorris,L., Erickson,J., Faro,S.,  
Ferreira,P., Fitzgerald,M., Gage,D., Galagan,J., Gardyna,S.,  
Graham,L., Grand-Pierre,N., Hafez,N., Hagopian,D., Hagos,B.,  
Hall,J., Horton,L., Hulme,W., Iliiev,I., Johnson,R., Jones,C.,  
Kamat,A., Karatas,A., Kellis,C., Landers,T., Levine,R.,  
Lindblad-Toh,K., Liu,G., Lui,A., Mabbitt,R., Maclean,C.,  
Macdonald,P., Major,J., Manning,J., Matthews,C., McCarthy,M.,  
Meldrim,J., Meneus,L., Mihova,T., Mlenga,V., Murphy,T., Naylor,J.,  
Nguyen,C., Nicol,R., Norbu,C., O'Connor,T., O'Donnell,P.,  
O'Neill,D., Oliver,J., Peterson,K., Phunkhang,P., Pierre,N.,  
Rachupka,A., Ramasamy,U., Raymond,C., Retta,R., Rise,C., Rogov,P.,  
Roman,J., Schauer,S., Schupback,R., Seaman,S., Severy,P., Smith,C.,  
Spencer,B., Stange-Thomann,N., Stojanovic,N., Stubbs,M.,  
Talamas,J., Tesfaye,S., Theodore,J., Topham,K., Travers,M.,  
Vassiliev,H., Venkataraman,V.S., Viel,R., Vo,A., Wilson,B., Wu,X.,  
Wyman,D., Young,G., Zainoun,J., Zembek,L., Zimmer,A. and Zody,M.  
Direct Submission  
Submitted (23-MAR-2003) Whitehead Institute/MIT Center for Genome  
Research, 320 Charles Street, Cambridge, MA 02141, USA  
On Mar 23, 2003 this sequence version replaced gi:28191504.  
All repeats were identified using RepeatMasker:  
Smit, A.F.A. & Green, P. (1996-1997)  
<http://ftp.genome.washington.edu/RM/RepeatMasker.html>

TITLE  
JOURNAL  
REFERENCE  
AUTHORS

FEATURES  
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/notes="assembly\_fragment"  
72037..76569  
/notes="assembly\_fragment"  
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/notes="assembly\_fragment"  
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Best Local Similarity 63.8%; Pred. No. 5.8e-31;  
Matches 307; Conservative 0; Mismatches 162; Indels 12; Gaps 3;

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DB 115085 AGAATATATTTCTTACCCAGGAGAAACATGATTGAGACCCAGCTAGAGTAGGATGA 115026  
QY 72 AGAATTCCTGCTGGTACTTTTCTTCCAGGAGCACTCTCTGATATTTTTTTTACA 131  
DB 115025 AGACTCCTTGCTGGTTCCTTTCTTTCTGGGAGAACAGTTCCTTAGACATTGATCTTACAA 114966

TITLE  
JOURNAL  
COMMENT

Center: Whitehead Institute/ MIT Center for Genome Research  
Center code: WIBR  
Web site: <http://www-seq.wi.mit.edu>  
Contact: [sequence\\_submissions@genome.wi.mit.edu](mailto:sequence_submissions@genome.wi.mit.edu)  
----- Project Information  
Center project name: L23317  
Center clone name: 82\_I\_24  
----- Summary Statistics  
Sequencing vector: Plasmid; n/a; 100% of reads  
Chemistry: Dye-terminator Big Dye; 100% of reads  
Assembly program: Phrap; version 0.960731  
Consensus quality: 205497 bases at least Q40  
Consensus quality: 206243 bases at least Q30  
Consensus quality: 206432 bases at least Q20

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 Db 114965 GCC-----AATGAATATTTTCGACCGTGGATTTCTATATTTCTCTTCTACAAAT 114914  
 QY 192 TCTAAACCTCTGACATTTGGTGAGACATTGAG---TACATTTTTTCCCATATCCCTACTT 248  
 Db 114913 GC AAAAAGTTGACCGTAGTGAGAAAGTTAAGCAAACTCTATTTCTATATGCTACTT 114854  
 QY 249 TTCAGAGAAATTTCTCTGCTGCTCCTCACTTAACATTTGCTGAGCGTCACTTTCTCTCC 308  
 Db 114853 TCGGAATGGCTTCTCTCCGCTGGTCCATTTGCTATCCCTGATGCGTCACTTTCTCTCC 114794  
 QY 309 TCATCTCTTTTCAGGCGCTCGAGAGCGAGGAGAGACAGAGGAGCTGTACTCAGAGCGG 368  
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 QY 428 AGGAGAGCGCTCAGAGGATTTCTGACAATATCTTTACCGGAGAGAGGCAAGTAGCTCA 487  
 Db 114673 ACCGAGAGGCTCAGAGGATTTCTGACAGCTTCTTTCAGAGCAGAGCAACTGTGCTCAA 114614  
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 Db 114613 A 114613

RESULT 5  
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 DEFINITION Rattus norvegicus clone CH230-9B12, WORKING DRAFT SEQUENCE, 4  
 ACCESSION AC125563  
 VERSION AC125563.4 GI:24817906  
 KEYWORDS HTG; HTGS PHASE1; HTGS DRAFT; HTGS\_FULLTOP.  
 SOURCE Rattus norvegicus (Norway rat)  
 ORGANISM Rattus norvegicus

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae;  
 Rattus.

1 (bases 1 to 275631)  
 Muzny, D. Marie, Metzker, M. Lee, Abramson, S., Adams, C., Alder, J.,  
 Allen, C., Allen, H., Albrooks, S., Amin, A., Anguiano, D.,  
 Bayalabechi, V., Ayodeji, A., Ayodeji, M., Baca, E., Baden, H.,  
 Baldwin, D., Bandaranaike, D., Barber, M., Barnstead, M., Benahmed, F.,  
 Biswal, K., Blair, J., Blankenburg, K., Blyth, P., Brown, M.,  
 Bryant, N., Buhay, C., Burch, P., Burrell, K., Calderon, E.,  
 Cardenas, V., Carter, K., Cavazos, I., Cesar, H., Chen, A.,  
 Chacko, J., Chavez, D., Chen, G., Chen, R., Chen, Y., Chen, Z., Chu, J.,  
 Cleveland, C., Cockrell, R., Cox, C., Coyle, M., Cree, A., D'Souza, L.,  
 Davila, M.L., Davis, C., Davy-Carroll, L., De Anda, C., Dederich, D.,  
 Delgado, O., Denson, S., Deramo, C., Ding, Y., Dinh, H., Divya, K.,  
 Draper, H., Dugan-Rocha, S., Dunn, A., Durbin, K., Duval, B., Eaves, K.,  
 Egan, A., Escotto, M., Eugene, C., Evans, C.A., Falls, T., Fan, G.,  
 Fernandez, S., Finley, M., Flagg, N., Forbes, L., Foster, M., Foster, P.,  
 Fraser, C.M., Gabisi, A., Ganta, R., Garcia, A., Garner, T., Garza, M.,  
 Gebregregis, E., Geer, K., Gill, R., Grady, M., Guerra, W., Guevara, W.,  
 Gunaratne, P., Haaland, W., Hamill, C., Hamilton, C., Hamilton, K.,  
 Harvey, J., Havlak, P., Hawes, A., Henderson, N., Hernandez, J.,  
 Hernandez, R., Hines, S., Hladik, S.L., Hodgson, A., Hogue, M.,  
 Hollins, B., Howells, S., Hulys, S., Hume, J., Idlebird, D., Jackson, A.,  
 Jackson, L., Jacob, L., Jiang, H., Johnson, B., Johnson, R., Jolivet, A.,  
 Karpach, S., Kelly, S., Kelly, S., Khan, Z., King, L., Kovar, C.,  
 Kowitz, J., Kraft, C.L., Lebow, H., Levan, J., Lewis, L., Li, Z., Liu, J.,  
 Liu, J., Liu, W., Liu, Y., London, P., Longacre, S., Lopez, J.,  
 Lorensuwa, L., Loulseghe, H., Lozano, R.J., Lu, X., Ma, J.,  
 Maheshwari, M., Mahindaratne, M., Mahmoud, M., Malloy, K., Mangum, A.,  
 Mangum, B., Mapua, P., Martin, K., Martin, R., Martinez, E.,  
 Mashiney, S., McLeod, M.P., McNeill, T.Z., Meenen, E.,  
 Milosavljevic, A., Miner, G., Minja, E., Montemayor, J., Moore, S.,

Morgan, M., Morris, K., Morris, S., Munidasa, M., Murphy, M., Nair, L.,  
 Nankervis, C., Neal, D., Newton, N., Nguyen, N., Norris, S.,  
 Nwankemelam, O., Okunou, G., Olarnpusagoon, A., Pal, S., Parks, K.,  
 Pasternak, S., Paul, H., Perez, A., Perez, L., Pfannkuch, C.,  
 Plapper, F., Poindexter, A., Popovic, D., Primus, E., Pu, L.,  
 Puazo, M., Quiroz, J., Rachlin, E., Reeves, K., Regier, M.A., Reigh, R.,  
 Reilly, B., Reilly, M., Ren, Y., Reuter, M., Richards, S., Riggs, F.,  
 Rives, C., Rodkey, T., Rojas, A., Rose, M., Rose, R., Ruiz, S.J.,  
 Sanders, W., Savary, G., Scherer, S., Scott, G., Shatsman, S., Shen, H.,  
 Shetty, J., Shvartsbeyn, A., Sisson, I., Sitter, C.D., Smales, D.,  
 Sneed, A., Sodergren, E., Song, X.-Z., Sorelle, R., Sosa, J.,  
 Steidle, M., Strong, R., Sutton, A., Svatek, A., Tabors, P., Taylor, C.,  
 Taylor, T., Thomas, N., Thomas, S., Tingey, A., Trejos, P., Usmani, K.,  
 Valas, R., Vera, V., Villalana, D., Waldron, L., Walker, B., Wang, J.,  
 Wang, Q., Wang, S., Warren, J., Warren, R., Wei, X., White, F.,  
 Williams, G., Willison, R., Wleczyk, R., Wood, H., Worley, K.,  
 Wright, D., Wright, R., Wu, J., Yakub, S., Yen, J., Yoon, L., Yoon, V.,  
 Yu, F., Zhang, J., Zhou, J., Zhou, X., Zhao, S., Dunn, D., von  
 Niederhausern, A., Weiss, R., Smith, D.R., Holt, R.A., Smith, H.O.,  
 Weinstein, G. and Gibbs, R.A.

TITLE  
 JOURNAL  
 REFERENCE  
 AUTHORS

TITLE  
 JOURNAL

REFERENCE  
 AUTHORS  
 TITLE  
 JOURNAL

COMMENT

Direct Submission  
 Unpublished  
 2 (bases 1 to 275631)  
 Worley, K.C.

Direct Submission  
 Submitted (29-JUN-2002) Human Genome Sequencing Center, Department  
 of Molecular and Human Genetics, Baylor College of Medicine, One  
 Baylor Plaza, Houston, TX 77030, USA  
 3 (bases 1 to 275631)  
 Rat Genome Sequencing Consortium.

Direct Submission  
 Submitted (09-NOV-2002) Human Genome Sequencing Center, Department  
 of Molecular and Human Genetics, Baylor College of Medicine, One  
 Baylor Plaza, Houston, TX 77030, USA

On Nov 9, 2002 this sequence version replaced gi:23096552  
 and whole genome shotgun sequencing reads assembled using Atlas  
 (http://www.hgsc.bcm.tmc.edu/projects/rat/). Each contig described  
 in the feature table below represents a scaffold in the Atlas  
 assembly (a 'contig-scaffold'). Within each contig-scaffold,  
 individual sequence contigs are ordered and oriented, and separated  
 by sized gaps filled with Ns to the estimated size. The sequence  
 may extend beyond the ends of the clone and there may be sequence  
 contigs within a contig-scaffold that consist entirely of whole  
 genome shotgun sequence reads. Both end sequences and whole genome  
 shotgun sequence only contigs will be indicated in the feature  
 table.

Center: Genome Center  
 Center: Baylor College of Medicine  
 Center code: BCM  
 Web site: http://www.hgsc.bcm.tmc.edu/  
 Contact: hgsc-help@bcm.tmc.edu  
 Project Information  
 Center project name: GDBK  
 Center clone name: CH230-9B12  
 Summary Statistics

Assembly program: Phrap; version 0.990329  
 Consensus quality: 228152 bases at least Q40  
 Consensus quality: 230310 bases at least Q30  
 Consensus quality: 231872 bases at least Q20  
 Estimated insert size: 235039; sum-of-contigs estimation  
 Quality coverage: 7x in Q20 bases; sum-of-contigs estimation

\* NOTE: Estimated insert size may differ from sequence length  
 \* (see http://www.hgsc.bcm.tmc.edu/docs/genbank\_draft\_data.html).  
 \* NOTE: This is a 'working draft' sequence. It currently  
 \* consists of 4 contigs. The true order of the pieces  
 \* is not known and their order in this sequence record is  
 \* arbitrary. Gaps between the contigs are represented as  
 \* runs of N, but the exact sizes of the gaps are unknown.  
 \* This record will be updated with the finished sequence  
 \* as soon as it is available and the accession number will  
 \* be preserved.

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* 1 192878: contig of 192878 bp in length
* 192878: gap of unknown length
* 192979 270395: contig of 77317 bp in length
* 270395: gap of unknown length
* 270396 272405: contig of 2010 bp in length
* 272405: gap of unknown length
* 272406 272505: gap of unknown length
* 272505: contig of 3126 bp in length.
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            complement(57750..59243)
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Best Local Similarity 64.0%; Pred. No. 1.3e-23;
Matches 309; Conservative 0; Mismatches 158; Indels 16; Gaps 6;
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QY 72 AGAATTCCTGCTGGTACTTTTCTCCAGGAAGCACTTCCTTGATATTTTTTTTTTACA 131
Db 142910 AGATCCTCTGCTGGTTCCTTTTCTGGGAGGCAAGTTCCTTT-AGACATGATTTTACA 142852
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Db 142851 AGCCACGAAATAGTCTCATATTTTTCAGAACCTGATTTCTATAGATCTCTCTACAA 142792
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Db 142791 AITCAGAACTTCACACATTTGTTAGAAATGTTGAGCAAA-----ACGTCCTTTTCTGTT 142739
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QY 310 CATCTCTTTACGGGCTGAGAGGCGAGAGG---AGACAGAGAGTGGTACTGCGAGAGC 366
Db 142678 C--CTTTCTAAAGGGCTTGAGAGGCGAGAGCGGCTCAGAGAGCGAGCGGATCTATAGCC 142621
QY 367 GCTCGTCTGATTCGCTGGAAGG--TCGTCAGTGGCTTATAAAGAGACCCCTACAGGCTTA 425
Db 142620 TGCCATCTGATTTGGCTTGGCGGGGGCTGGCTGGGCTTATAAAGAGACCCCTACAGCTTG 142561
QY 426 GCAGGAGAGCGCTCAGAGGATTTGCAATAATCTTTACCGGAGAGAGGAGGCAAGTACGCT 485
Db 142560 AGAGGAGAGGCTCAGAGGATTTGCAAGCTTCTTTGAGAGGAGGAGCAAGTGTGCTCA 142501
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Db 142500 AAA 142498
AK093959 2599 bp mRNA linear PRI 15-JUL-2002
LOCUS
RESULT 6
AK093959
LOCUS

```

Homo sapiens cDNA FLJ36640 fis, clone TRACH2019151, moderately similar to REGULATOR OF G-PROTEIN SIGNALING 4.

AK093959 1 GI:21752824  
oligo capping; fis (full insert sequence).  
Homo sapiens (human)  
Homo sapiens  
Rukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

1 Suzuki, O., Sasaki, N., Aotsuka, S., Shoji, T., Ichihara, T., Shiohata, N., Matsumoto, K., Hirano, M., Sano, S., Nomura, K., Yoshikawa, Y., Matsumura, Y., Moriya, S., Chiba, E., Moniyama, H., Onogawa, S., Kaeriyama, S., Satoh, N., Matsumura, H., Takahashi, E., Kataoka, R., Kuga, N., Kuroda, A., Satoh, I., Kamata, K., Takami, S., Terashima, Y., Watanabe, M., Sugiyama, T., Irie, R., Otsuki, T., Sato, H., Ota, T., Wakamatsu, A., Ishii, S., Yamamoto, J., Isono, Y., Kawai-Hio, Y., Saito, K., Nishikawa, T., Kimura, K., Yamashita, H., Matsuo, K., Nakamura, Y., Sekine, M., Kikuchi, H., Kanda, K., Magatsuma, M., Murakawa, K., Kanehori, K., Takahashi-Fujii, A., Oshima, A., Sugiyama, A., Kawakami, B., Suzuki, Y., Sugano, S., Nagahara, K., Masubo, Y., Nagai, K. and Isogai, T.

NEDO human cDNA sequencing project  
Unpublished  
2 (bases 1 to 2599)  
Isogai, T. and Yamamoto, J.  
Direct Submission  
Submitted (04-JUL-2002) Takao Isogai, FLJ Project (HRI Team); 2-6-7 Kazusa-Kamatari, Kisarazu, Chiba 292-0812, Japan (E-mail: genomics@hri.co.jp, Tel: 81-438-52-3975, Fax: 81-438-52-3986)  
NEDO human cDNA sequencing project supported by Ministry of Economy, Trade and Industry of Japan; cDNA full insert sequencing: Research Association for Biotechnology (RAB); cDNA library construction: Helix Research Institute (HRI) (supported by Japan Key Technology Center etc.); 5'- & 3'-end one pass sequencing: RAB, HRI, and Biotechnology Center, National Institute of Technology and Evaluation; clone selection for full insert sequencing: HRI and RAB; annotation: HRI and RAB.

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Matches 70; Conservative 0; Mismatches 1; Indels 0; Gaps 0;  
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Db 1 AAGACGCTCAGAGGATTTCTGCAATATCTTTTACCGAGAGAGGCGGAGTACGCTCAAG 60  
QY 491 CCGAAGCCACA 501  
Db 61 CCGAAGCCACA 71

RESULT 7  
AK096204  
LOCUS  
DEFINITION Homo sapiens cDNA FLJ38885 fis, clone MESAN2017417, moderately similar to REGULATOR OF G-PROTEIN SIGNALING 4.  
ACCESSION AK096204  
VERSION AK096204.1 GI:21755635  
KEYWORDS oligo capping; fis (full insert sequence).  
SOURCE Homo sapiens (human)  
ORGANISM Homo sapiens

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

1 Tanigami, A., Fujiwara, T., Shibahara, T., Goto, Y., Hirao, M., Shimizu, F., Wakebe, H., Ono, T., Hishigaki, H., Matanabe, T., Ozaki, K., Sugiyama, T., Irie, R., Otsuki, T., Sato, H., Wakamatsu, A., Ishii, S., Yamamoto, J., Isono, Y., Kawai-Hio, Y., Saito, K., Nishikawa, T., Kimura, K., Yamashita, H., Matsuo, K., Nakamura, Y., Sekine, M., Kikuchi, H., Kanda, K., Wagatsuma, M., Murakawa, K., Kanehori, K., Takahashi-Fujii, A., Oshima, A., Sugiyama, A., Kawakami, B., Suzuki, Y., Sugano, S., Negahari, K., Masuho, Y., Nagai, K. and Isogai, T. NEDO human cDNA sequencing project

Unpublished  
2 (bases 1 to 3769)  
Isogai, T. and Yamamoto, J.  
Direct Submission  
Submitted (04-JUL-2002) Takao Isogai, FLJ Project (HRI Team); 2-6-7 Kazusa-Kamatari, Kisarazu, Chiba 292-0812, Japan (E-mail: genomics@ri.co.jp, Tel: 81-438-52-3975, Fax: 81-438-52-3986) NEDO human cDNA sequencing project supported by Ministry of Economy, Trade and Industry of Japan; cDNA full insert sequencing: Research Association for Biotechnology (RAB); cDNA library construction: Helix Research Institute (HRI) (supported by Japan Key Technology Center etc.); 5'- & 3'-end one pass sequencing: RAB, HRI, and Biotechnology Center, National Institute of Technology and Evaluation; clone selection for full insert sequencing: HRI and RAB; annotation: HRI and RAB.

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Best Local Similarity 100.0%; Pred. No. 1.7e-06;  
Matches 69; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

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QY 493 GAAGCCACA 501  
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DB 61 GAAGCCACA 69

RESULT 8  
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LOCUS  
DEFINITION  
Rattus norvegicus clone CH230-158A12, WORKING DRAFT SEQUENCE, 3 unordered pieces.  
AC111539  
AC111539.5 GI:30579258  
HTG; HTGS\_PHASE1; HTGS\_DRAFT; HTGS\_FULLTOP.  
Rattus norvegicus (Norway rat)  
Rattus norvegicus  
ORGANISM  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.

REFERENCE  
1 (bases 1 to 234560)  
Muzny, D., Marie, J., Metzker, M., Lee, J., Abramson, S., Adams, C., Alder, J., Allen, C., Allen, H., Alsbrooks, S., Amin, A., Anguiano, D., Anyalebechi, V., Aoyagi, A., Ayodeji, M., Baca, B., Baden, H., Baldwin, D., Bandaranaike, D., Barber, M., Barnstead, M., Benahmed, F., Biswal, K., Blair, J., Blankenburg, K., Blyth, P., Brown, M., Bryant, N., Buhay, C., Burch, P., Burrell, K., Calderon, E.,

Cardenas, V., Carter, K., Cavazos, I., Ceasar, H., Center, A., Chacko, J., Chavez, D., Chen, G., Chen, R., Chen, Y., Chen, Z., Chu, J., Cleveland, C., Cockrell, R., Cox, C., Coyle, M., Cree, A., D'Souza, L., Davila, M., Davis, C., Davy-Carroll, L., De Anda, C., Dederich, D., Delgado, O., Denison, S., Deramo, C., Ding, Y., Dinh, H., Diyva, K., Draper, H., Dugan-Rocha, S., Dunn, A., Durbin, K., Duval, B., Evans, K., Egan, A., Escotto, M., Eugene, C., Evans, C. A., Falls, T., Fan, G., Fernandez, S., Finley, M., Flagg, N., Forbes, L., Foster, M., Foster, P., Fraser, C. M., Gabisi, A., Ganta, R., Garcia, A., Garner, T., Garza, M., Gbrageorgis, E., Geer, K., Gill, R., Grady, M., Guerra, W., Guevara, W., Gunaratne, P., Haaland, W., Hamil, C., Hamilton, C., Hamilton, K., Harvey, Y., Haviak, P., Hawes, A., Henderson, N., Hernandez, J., Hernandez, R., Hines, S., Hladun, S. L., Hodgson, A., Hogues, M., Hollins, B., Howells, S., Hulyk, S., Hume, J., Idlebird, D., Jackson, A., Jackson, L., Jacob, L., Jiang, H., Johnson, B., Johnson, R., Jolivet, A., Karpathy, C., Kelly, S., Kelly, S., Khan, Z., King, L., Kovar, C., Kowis, C., Kraft, C. L., Lebow, H., Levan, J., Lewis, L., Li, Z., Liu, J., Liu, J., Liu, W., Liu, Y., London, P., Longacre, S., Lopez, J., Lorenshewa, L., Louised, H., Lozano, R. J., Lu, X., Ma, J., Maheshwari, M., Mahindartne, M., Mahmoud, M., Malloy, K., Mangum, A., Mangum, B., Mapua, P., Martin, K., Martin, R., Martinez, E., Mawhinney, S., McLeod, M. P., McNeill, T. Z., Meenen, E., Milosavljevic, A., Miner, G., Minja, E., Montemayor, J., Moore, S., Morgan, M., Morris, K., Morris, S., Munidasa, M., Murphy, M., Nair, L., Nankervis, C., Neal, D., Newton, N., Nguyen, N., Norris, S., O'Brien, S., Pasternak, S., Paul, H., Perez, A., Perez, L., Pfannkuch, C., Plopper, F., Poindexter, A., Popovic, D., Primus, E., Pu, L., L., Puazo, M., Quiroz, J., Rachlin, E., Reeves, K., Regier, M. A., Reigh, R., Reilly, B., Reilly, M., Ren, Y., Reuter, M., Richards, S., Riggs, P., Rives, C., Rodkey, T., Rojas, A., Rose, M., Rose, R., Ruiz, S. J., Sanders, W., Savery, G., Scherer, S., Scott, G., Shatsman, S., Shen, H., Shetty, J., Shvartsbeyn, A., Sisson, I., Sittler, C. D., Smajda, D., Sneed, A., Sodergren, E., Song, X.-Z., Sorelle, R., Sosa, J., Steimle, M., Strong, R., Sutton, A., Svatek, A., Taber, P., Taylor, C., Taylor, T., Thomas, N., Thomas, S., Tingey, A., Trejos, Z., Usmani, K., Valas, R., Vera, V., Villaseña, D., Waldron, L., Walker, B., Wang, J., Wang, Q., Wang, S., Warren, J., Warren, R., Wei, X., White, F., Williams, G., Willson, R., Wlezyk, R., Wooden, H., Worley, K., Wright, D., Wright, R., Wu, J., Yakub, S., Yen, J., Yoon, L., Yoon, V., Yu, F., Zhang, J., Zhou, X., Zhao, S., Dunn, D., von Niederhausern, A., Weiss, R., Smith, D. R., Holt, R. A., Smith, H. O., Weinstock, G. and Gibbs, R. A.

Direct Submission  
Unpublished  
2 (bases 1 to 234560)  
Worley, K. C.  
Direct Submission  
Submitted (19-FEB-2002) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
3 (bases 1 to 234560)  
Rat Genome Sequencing Consortium.  
Direct Submission  
Submitted (13-MAY-2003) Human Genome Sequencing Center, Department of Molecular and Human Genetics, Baylor College of Medicine, One Baylor Plaza, Houston, TX 77030, USA  
On May 13, 2003 this sequence version replaced gi:25007559.  
The sequence in this assembly is a combination of BAC based reads and whole genome shotgun sequencing reads assembled using Atlas (<http://www.hgsc.bcm.tmc.edu/projects/rat/>). Each contig described in the feature table below represents a scaffold in the Atlas assembly (a 'contig-scaffold'). Within each contig-scaffold, individual sequence contigs are ordered and oriented, and separated by sized gaps filled with Ns to the estimated size. The sequence may extend beyond the ends of the clone and there may be sequence contigs within a contig-scaffold that consist entirely of whole genome shotgun sequence reads. Both end sequences and whole genome shotgun sequence only contigs will be indicated in the feature table.

----- Genome Center of Medicine  
Center: Baylor College of Medicine  
Center code: BCM





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repeat_region 102..187
repeat_region /rpt_family="AT_rich"
repeat_region 256..285
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repeat_region 656..687
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repeat_region /complement(865..960)
repeat_region /rpt_family="MIR"
repeat_region complement(971..1278)
repeat_region /rpt_family="AluY"
repeat_region /complement(1728..2029)
repeat_region /rpt_family="AluSg"
repeat_region 2425..2653
repeat_region /rpt_family="L1M4"
repeat_region 3273..3514
repeat_region /rpt_family="MIR"
repeat_region 3780..3833
repeat_region /rpt_family="AT_rich"
repeat_region /complement(3834..3939)
repeat_region /rpt_family="L1PA5"
repeat_region /complement(4079..4371)
repeat_region /rpt_family="AluDb"
repeat_region /complement(5881..6125)
repeat_region /rpt_family="MLT1B"
repeat_region /complement(6126..9075)
repeat_region /rpt_family="L1PA13"
repeat_region /complement(9076..9252)
repeat_region /rpt_family="MLT1B"
repeat_region 10440..10582
repeat_region /rpt_family="HAL1b"
repeat_region /complement(10583..10875)
repeat_region /rpt_family="AluSg"
repeat_region 10876..10966
repeat_region /rpt_family="HAL1b"
repeat_region /complement(10995..11175)
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repeat_region 11375..12064
repeat_region /rpt_family="L2"
repeat_region 12273..12344
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repeat_region 12350..12394
repeat_region /rpt_family="AT_rich"
repeat_region 12395..12876
repeat_region /rpt_family="L2"
repeat_region 12996..13055
repeat_region /rpt_family="TAGA)n"
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repeat_region /rpt_family="L1MB4A"
repeat_region 13654..13760
repeat_region /rpt_family="AluSg/x"
repeat_region 13869..13908
repeat_region /rpt_family="AT_rich"
repeat_region 14076..14144
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repeat_region 14333..14353
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repeat_region /complement(14675..15120)
repeat_region /rpt_family="L3"
repeat_region complement(15749..18697)
repeat_region /rpt_family="L1MA4"
repeat_region /complement(18698..18997)
repeat_region /rpt_family="AluSp"
repeat_region /complement(18998..20399)
repeat_region /rpt_family="L1MA4"
repeat_region 20400..20707
repeat_region /rpt_family="AluY"
repeat_region /complement(20708..21184)
repeat_region /rpt_family="L1MA4"
repeat_region /complement(21178..22061)
repeat_region /rpt_family="L1MA6"
repeat_region /complement(22062..24224)
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unsure 25698..25704
repeat_region /note="<30 qual SNGL region"
repeat_region 25996..26034
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repeat_region 28711..28766
repeat_region /rpt_family="MER112"
repeat_region 28767..29060
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repeat_region /rpt_family="MER112"
repeat_region /complement(29175..29326)
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repeat_region /complement(29534..29652)
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repeat_region 30421..30448
repeat_region /rpt_family="AT_rich"
repeat_region 30459..30752
repeat_region /rpt_family="AluSx"
repeat_region 30932..31224
repeat_region /rpt_family="AluJo"
repeat_region 31264..32303
repeat_region /rpt_family="L1PAL0"
repeat_region 33754..33774
repeat_region /rpt_family="AT_rich"
repeat_region 35349..35651
repeat_region /rpt_family="AluSx"
repeat_region /complement(36456..36759)
repeat_region /rpt_family="AluSx"
repeat_region /complement(38255..38613)
repeat_region /rpt_family="MLT1C"
repeat_region /complement(39708..39864)
repeat_region /rpt_family="L1P3"
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Best Local Similarity 60.9%; Pred. No. 0.32;
Matches 78; Conservative 0; Mismatches 50; Indels 0; Gaps 0;

QY 106 AACTTCCTTGATATTTTTTTTACAGGCATATGAATAAACTATATTTGCACATG 165
Db 44705 AGCTTCCTGGCTTTTACATATAGTCTTATATTTACAGTATATTTAACCTCTTTG 44764

QY 166 TACACTTTTTTCTTTCTAGAAATCTAAACCTCTGACATTTGGTGAGACATTTAGTA 225
Db 44765 TTGACATGTTTACCTTTCCAATCAATCTTCACTCTGATATTTGGTTTCTTATCTT 44824

QY 226 CATTTTTT 233
Db 44825 GATTTTAT 44832
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RESULT 10
LOCUS I66494
DEFINITION Sequence 14 from patent US 5670367.
ACCESSION I66494
VERSION I66494.1 GI:2724471
KEYWORDS
SOURCE Unknown.
ORGANISM Unknown.
REFERENCE
1 (bases 1 to 7218)
AUTHORS Dörner, F., Scheiflinger, F. and Falkner, F. Gunter.
TITLE Recombinant fowlpox virus
JOURNAL Patent: US 5670367-A 14 23-SEP-1997;
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FEATURES	Location/Qualifiers
source	1. 7218
BASE COUNT	1944 a 1491 c 1486 g 1929 t 368 others
ORIGIN	
Query Match	9.5%; Score 47.6; DB 6; Length 7218;
Best Local Similarity	3.1%; Pred. No. 0.35;
Matches	8; Conservative 158; Mismatches 92; Indels 0; Gaps 0;
QY	62 AGTAAGCTCCAGAAATTCCTGCTGTACTTTCCTTCAGGAGCAACTCTTCGTATATTT 121
Db	1053 AGGAGCTCGCATYY 1112
QY	122 TTTTATACAGGCATATGAATAAACTATATTTTCGAGCATGTACACATTTTTCCTT 181
Db	1113 YY 1172
QY	182 TTCTAGAAATCTAAACCTCTGACATTTGGTGAGACATTCAGTACATTTTTCCTATC 241
Db	1173 YY 1232
QY	242 CCTACTTTTCAGAGGATTTTCTCTGCTCGTTCTACCTTAACTGCTGATCGTCAGTCTT 301
Db	1233 YY 1292
QY	302 TTCTTCTCATCTCTTTC 319
Db	1293 YYYYYYYYYYYYYYYYYY 1310
RESULT 11	
AC095234	
LOCUS	307979 bp DNA linear HTG 26-SEP-2002
DEFINITION	Rattus norvegicus clone CH230-10H10, *** SEQUENCING IN PROGRESS
ACCESSION	AC095234
VERSION	AC095234.4 GI:22772995
KEYWORDS	HTG; HTGS PHASE1; HTGS DRAFT; HTGS_ENRICHED.
SOURCE	Rattus norvegicus
ORGANISM	Rattus norvegicus (Norway rat)
Eukaryote; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Rattus.	
REFERENCE	1 (bases 1 to 307979)
AUTHORS	Muzny,D,Marle, Metzker,M, Lee, Abramson,S, Adams,C, Alder,J, Allen,C, Allen,H, Alsbrooks,S, Amin,A, Anguano,D, Anyatebechi,V, Aoyagi,A, Ayodeji,M, Baca,E, Baden,H, Baldwin,D, Bandaranaike,D, Barber,M, Barnstead,M, Benahmed,F, Biswal,N, Blair,J, Blankenburg,K, Blyth,P, Brown,M, Bryant,N, Buahy,C, Burch,P, Burrell,K, Calderon,E, Cardenas,V, Carter,K, Cavazos,I, Ceasar,H, Center,A, Chacko,J, Chavez,D, Chen,G, Chen,R, Chen,Y, Chen,Z, Chu,J, Cleveland,C, Cockrell,R, Cox,C, Coyle,M, Cree,A, D'Souza,L, Davila,M,L, Davis,C, Davy-Carroll,L, De Anda,C, Dederich,D, Delgado,O, Denson,S, Deramo,C, Ding,Y, Dinh,H, Divya,K, Draper,H, Dugan-Rocha,S, Dunn,A, Durbin,K, Duval,B, Eaves,K, Egan,A, Escotto,M, Eugene,C, Evans,C,A, Falls,T, Fan,G, Fernandez,S, Finley,M, Flagg,N, Forbes,L, Foster,M, Foster,P, Fraser,C,M, Gabisi,A, Ganta,R, Garcia,A, Garner,T, Garza,M, Gatregeorgis,E, Geer,K, Gill,R, Grady,M, Guerra,W, Guevara,W, Gunaratne,P, Haaland,W, Hamil,C, Hamilton,C, Hamilton,K, Harvey,Y, Havlak,P, Hawes,A, Henderson,N, Hernandez,J, Hernandez,R, Hines,S, Hladun,S,L, Hodgson,A, Hogues,M, Hollins,B, Howells,S, Hulyk,S, Hume,J, Idlebird,D, Jackson,A, Jackson,L, Jacob,L, Jiang,H, Johnson,B, Johnson,R, Jolivet,A, Karpathy,S, Kelly,S, Kelly,S, Khan,Z, King,L, Kovar,C, Kowis,C, Kraft,C,L, Lebow,H, Levan,J, Lewis,L, Li,Z, Liu,J, Liu,J, Liu,W, Liu,Y, London,P, Longacre,S, Lopez,J, Lorensuewa,L, Loulsegod,H, Lozada,R,J, Lu,X, Ma,J, Maheshwari,M, Mahindartne,M, Mahmoud,M, Mallory,K, Mangum,A, Mangum,B, Mapua,P, Martin,K, Martin,R, Martinez,E, Mawhiney,S, McLeod,M,P, McNeill,T,Z, Meenen,E,

\* 231416 231515: gap of unknown length  
 \* 231516 237693: contig of 6178 bp in length  
 \* 237694 237793: gap of unknown length  
 \* 237794 287892: contig of 50099 bp in length  
 \* 287893 287992: gap of unknown length  
 \* 287993 297230: contig of 9238 bp in length  
 \* 297231 297330: gap of unknown length  
 \* 297331 301900: contig of 4570 bp in length  
 \* 301901 302000: gap of unknown length  
 \* 302001 303486: contig of 1486 bp in length  
 \* 303487 303586: gap of unknown length  
 \* 303587 304819: contig of 1233 bp in length  
 \* 304820 304919: gap of unknown length  
 \* 304920 307979: contig of 3060 bp in length.

## FEATURES

Location/Qualifiers

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/organism="Rattus norvegicus"

/mol\_type="genomic DNA"

/db\_xref="taxon:10116"

/clone="CH230-10H10"

3381..5706

/note="wgs\_end\_extension"

clone\_end:Sp6"

10057..10967

/note="clone boundary"

clone\_end:Sp6"

site:EcoRI

end\_sequence:BH305632"

21440..23858

/note="wgs\_contig"

30185..33987

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51818..53151

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80233..82290

/note="wgs\_contig"

187059..187447

/note="clone boundary"

clone\_end:T7"

site:EcoRI

end\_sequence:BH305630"

231516..233344

/note="wgs\_end\_extension"

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234226..235378

/note="wgs\_end\_extension"

clone\_end:T7"

236591..237693

/note="wgs\_end\_extension"

clone\_end:T7"

298726..301900

/note="wgs\_end\_extension"

clone\_end:T7"

BASE COUNT 67479 a 47083 c 45283 g 66098 t 82036 others

## ORIGIN

Query Match 9.1%; Score 45.6; DB 2; Length 307979;

Best Local Similarity 50.4%; Pred. No. 1.3; Indels 2; Gaps 1;

Matches 138; Conservative 0; Mismatches 134; Indels 134; Gaps 1;

QY 10 TAATATGATCTCTTAATCCAAAGAGGAAGGCAATGGAGTCAGCTCCTAAGTAAGCT 69

Db 132895 TTATTGTGGCCACTTGCTGAAATCAAAAGGTGGTCAGTCTCTTCACTTTGTAACT 132954

QY 70 CCAGATTCCTGCT--GGTACTTTCTTCCTCAGGAGCACTTCCTTGATATTTTTTT 127

Db 132955 CCCCTTCCTCAATCTCAGGACCTAAACATATAGTAGTGAAGTGGCATTGATATCTCTGG 133014

QY 128 TACAGGCATATGATAAAATATATTTTGCAGCAATGTACACTTTTTCCTTTCTAG 187

Db 133015 TTCCCTATATTTTAAATTTTAAATTTTATAGATATATTTCTTTACTTACATTTCAA 133074

QY 188 AAATCTAAACCTCTGACATTTGGTGAGACATTTAGTACATATTTTCCCATCTCCTACT 247

Db 133075 CATTATTCCTCTCCCAATTTCTGTATATAAGCACCCATTCCTTCCCATCCCTCCC 133134

QY 248 TTTCAGAAGGATTTTCTCTGCTCGTTTCACCTTAAC 281

Db 133135 CTATAGAGGTATCCCTTATATACATCAGTTTAC 133168

## RESULT 12

AX347195

LOCUS

AX347195

DEFINITION

Sequence 2266 from Patent WO0200928.

ACCESSION

AX347195

VERSION

AX347195.1

KEYWORDS

synthetic construct

SOURCE

synthetic construct

ORGANISM

artificial sequences.

REFERENCE

1

AUTHORS

Olek,A., Piepenbrock,C. and Berlin,K.

TITLE

Diagnosis of diseases associated with the immune system

JOURNAL

Patent: WO 0200928-A 2266 03-JAN-2002;

EpiGenomics AG (DE)

FEATURES

Location/Qualifiers

1..10020

/organism="synthetic construct"

/mol\_type="genomic DNA"

/db\_xref="taxon:32630"

/note="chemically treated genomic DNA (Homo sapiens)"

BASE COUNT

3111 a 92 c 2113 g 4704 t

ORIGIN

Query Match

Best Local Similarity

9.1%; Score 45.4; DB 6; Length 10020;

Matches 132; Conservative

0; Mismatches 126; Indels

1; Gaps

1;

QY 90 TTTCCTCCAGGAAGCACTTCTGTATATTTTTTTTACAGGCATATGAATAAAACT 149

Db 1329 TTTGTTTTAGGATGAATTAGTTCGAGTTTTTTTTTTTATGATATATAAGGAAT 1388

QY 150 ATATTTTGCAGCATTTGACACTTTTTTCTCTAGAAATCTAAACCTCTGACATTCG 209

Db 1389 TTTTATTTGGTGTATGTTAGTTTTTTTATTTTGTGAATATAAATTTTGTGTTTG 1448

QY 210 GTGAGACATTCAGTACATTTTTTCCCATATCCCTACTTTTCAGAGGATTTTCTCTGCT 269

Db 1449 ATTTA-AGAAAGTTACGGTTTTTTTATTTTGTGTTTGTGTTTTTTTATTT 1507

QY 270 CGTTCACCTTAACATTCCTGATCGTCAGTCCTTTCTCTCATCTTTCAGGGGCTGGA 329

Db 1508 TTATTTTTTTTATTTTTTTTTTTTTTTTATATAAAGCTTTTAAATATGGA 1567

QY 330 GAGGCAGAGGGAGACAGAG 348

Db 1568 GAGAGAGAGAGAGAGAG 1586

## RESULT 13

AC116367/c

LOCUS

AC116367

DEFINITION

Oryza sativa (japonica cultivar-group) chromosome 11 clone

OSUNBa0059H21, \*\*\* SEQUENCING IN PROGRESS \*\*\*; 5 ordered pieces.

ACCESSION

AC116367

VERSION

AC116367.9

KEYWORDS

HTG; HTGS PHASE2.

SOURCE

Oryza sativa (japonica cultivar-group)

ORGANISM

Eukaryota; Viridiplantae; Streptophyta; Embryophyta; Tracheophyta;

Spermatophyta; Magnoliophyta; Liliopsida; Poales; Poaceae;

Ehrhartoideae; Oryzaceae; Oryza.

REFERENCE

1 (bases 1 to 134971)

AUTHORS

Buell,C.R., Yuan,Q., Ouyang,S., Liu,J., Gansberger,K., Jones,K.M.,

Overton II,L., Tsitrin,I., Kim,M., Bera,J., Jin,S., Fadrosch,D.W.,

SEQUENCE.  
AC022764 GI:11038547  
AC022764.4 HTGS PHASE2; HTGS\_DRAFT.  
HTG; HTGS PHASE2; HTGS\_DRAFT.  
Homo sapiens (human)  
Homo sapiens  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.  
1 (bases 1 to 198464)  
Birren,B., Linton,L., Nusbaum,C. and Lander,E.  
Homo sapiens chromosome 17, clone RP11-310M15  
Unpublished  
2 (bases 1 to 198464)  
Birren,B., Linton,L., Nusbaum,C., Lander,E., Abraham,H., Allen,N.,  
Anderson,S., Baldwin,J., Barna,N., Beckerly,R., Beda,F.,  
Boguslavsky,L., Bouckigalter,B., Brown,A., Burkett,G., Castle,A.,  
Choepel,Y., Collangelo,M., Collins,S., Collymore,A., Cooke,P.,  
DeAillano,K., Dewar,K., Domino,M., Doyle,M., Fenesor,J.,  
Ferreira,P., FitzHugh,W., Forrest,C., Gage,D., Galagan,J.,  
Gardyna,S., Grant,G., Hegos,B., Heaford,A., Horton,L.,  
Howland,J.C., Johnson,R., Jones,C., Kann,L., Karatas,A., Klein,J.,  
Landers,T., Lehotsky,J., Levine,R., Liu,C., Liu,G., Locke,K.,  
Macdonald,P., Marquis,N., McEwan,P., McGurk,A., McKernan,K.,  
McPheters,R., Meldrum,J., Meneus,L., Morrow,J., Naylor,J.,  
Norman,C.H., O'Connor,T., O'Donnell,P., Olivar,T.M., Peterson,K.,  
Pierre,N., Pisan,C., Pollava,V., Raymond,C., Riley,R., Rothman,D.,  
Roy,A., Santos,R., Severy,P., Spencer,B., Stange-Thomann,N.,  
Stojanovic,N., Subramanian,A., Talamas,J., Tesfaye,S., Theodore,J.,  
Tirrell,A., Vassiliev,H., Viel,R., Vo,A., Wu,X., Wyman,D., Ye,W.J.,  
Zimmer,A. and Zody,M.  
Direct Submission  
Submitted (06-FEB-2000) Whitehead Institute/MIT Center for Genome  
Research, 320 Charles Street, Cambridge, MA 02141, USA  
On Oct 30, 2000 this sequence version replaced gi:7429221.  
All repeats were identified using RepeatMasker:  
Smit, A.F.A. & Green, P. (1996-1997)  
<http://ftp.genome.washington.edu/RM/RepeatMasker.html>  
----- Genome Center  
Center: Whitehead Institute/ MIT Center for Genome Research  
Center code: WIBR  
Web site: <http://www-seq.wi.mit.edu>  
Contact: [sequence\\_submissions@genome.wi.mit.edu](mailto:sequence_submissions@genome.wi.mit.edu)  
----- Project Information  
Center project name: LS848  
Center clone name: 310.M.15  
----- Summary Statistics  
Sequencing vector: M13; M7815; 32% of reads  
Sequencing method: Plasmid; N/A; 68% of reads  
Chemistry: Dye-terminator Big Dye; 100% of reads  
Assembly program: Phrap; version 0.960731  
Consensus quality: 198356 bases at least Q40  
Consensus quality: 198459 bases at least Q30  
Consensus quality: 198464 bases at least Q20  
Insert size: 210000; agarose-fp  
Insert size: 198464; sum-of-contigs  
Quality coverage: 9.2 in Q20 bases; agarose-fp  
Quality coverage: 9.7 in Q20 b.  
\* NOTE: This is a 'working draft' sequence. It currently  
\* consists of 1 contigs. Gaps between the contigs  
\* are represented as runs of N. The order of the pieces  
\* is believed to be correct as given, however the sizes  
\* of the gaps between them are based on estimates that have  
\* provided by the submittor.  
\* This sequence will be replaced  
\* by the finished sequence as soon as it is available and  
\* the accession number will be preserved.  
\* 1 198464: contig of 198464 bp in length.  
assembly\_location.  
Location/Qualifiers  
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/db\_xref="taxon:9606"

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/map="17"
/clone="RP11-310M15"
/clone_lib="RPC1-11 Human Male BAC"
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ORIGIN
Query Match      9.0%; Score 45.2; DB 2; Length 198464;
Best Local Similarity 50.0%; Pred. No. 1.6; Indels 0; Gaps 0;
Matches 113; Conservative 0; Mismatches 113;
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Db      55513 TCAATCACTTCTACGATCTGTAATATTTTGTCTCTCTGAGTCATATATATATATATATAT 55572
QY      120  TTTTCTTTTACAGGGCATATGAATAAAACTATATTTTGAGCATTGTACACATTTTTTCTCC 179
Db      55573 ATTTTTTTCCTTCGTATATATTAATGTGCATTTTTTATTTCAATTTAAATTTATTTGT 55632
QY      180  TTTTCTAGAAATCTTAAACCTCTGACATTTGGTGAGACATTTGAGTACATTTTTTCCCAT 239
Db      55633 ATGACCTGTACATAAAATTTTATTAGAATTTTAAATATATCTCTATAAGTATTTTCCCTC 55692
QY      240  TCCTACTTTTCAGAAAGGATTTCTCTGCTGTTCACTTAAACATTC 285
Db      55693 TGGTACTAATGTTAATTTTACTTTTGTCTATTTTCTTTAATATGG 55738

RESULT 15
AL451076/c
LOCUS
DEFINITION
Mouse DNA sequence from clone RP23-43020 on chromosome X, complete
AL451076      203581 bp      DNA      linear      ROD 29-JUN-2002

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AL451076/C	LOCUS	203581 bp	DNA	linear	ROD 29-JUN-2002
	DEFINITION		Mouse DNA sequence from clone RP23-43020	on chromosome X, complete	

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ACCESSION  
AL451076.14 GI:20068432  
VERSION  
HTG.  
KEYWORDS  
SOURCE  
Mus musculus (house mouse)  
ORGANISM  
Mus musculus  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.  
REFERENCE  
1 (bases 1 to 203581)  
AUTHORS  
Phillimore, B.  
TITLE  
Direct Submission  
JOURNAL  
Submitted (29-JUN-2002) Wellcome Trust Sanger Institute, Hinxton,  
Cambridgeshire, CB10 1SA, UK. E-mail enquiries:  
humquery@sanger.ac.uk Clone requests: clonerequest@sanger.ac.uk  
On Apr 7, 2002 this sequence version replaced gi:14161186.  
During sequence assembly data is compared from overlapping clones.  
Where differences are found these are annotated as variations  
together with a note of the overlapping clone name. Note that the  
variation annotation may not be found in the sequence submission  
corresponding to the overlapping clone, as we submit sequences with  
only a small overlap as described above.  
This sequence was finished as follows unless otherwise noted: all  
regions were either double-stranded or sequenced with an alternate  
chemistry or covered by high quality data (i.e., phred quality >=  
30); an attempt was made to resolve all sequencing problems, such  
as compressions and repeats; all regions were covered by at least  
one plasmid subclone or more than one M13 subclone; and the  
assembly was confirmed by restriction digest. The following  
abbreviations are used to associate primary accession numbers given  
in the feature table with their source databases: Em: EMBL; Sw:,  
SWISSPROT; Tr:, TREMBL; Wp:, WORMPEP; Information on the WORMPEP  
database can be found at  
http://www.sanger.ac.uk/Projects/c\_elegans/wormpep RP23-43020 is  
from the RPC1-23 Mouse PAC Library  
constructed by the group of Pieter de Jong.  
For further details see http://www.chori.org/bacpac/home.htm  
VECTOR: pBACe3.6  
----- Genome Center  
Center: UK Medical Research Council  
Center code: UK-MRC  
Web site: http://mrcseq.har.mrc.ac.uk

GenCore version 5.1.6  
Copyright (c) 1993 - 2003 Compugen Ltd.

OM nucleic - nucleic search, using sw model

Run on: November 7, 2003, 01:54:16 ; Search time 1962.4 Seconds  
(without alignments)  
10444.222 Million cell updates/sec

Title: US-09-939-209a-3\_COPY\_15000\_15500  
Perfect score: 501  
Sequence: 1 tggcagagaactctctgat.....tcaacctgatgagaagat 501

Scoring table: IDENTITY NUC  
Gapop 10.0 , Gapext 1.0

Searched: 2888711 reqs, 20454813386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database :

GenEmbl.\*

- 1: gb.ba.\*
- 2: gb.htg.\*
- 3: gb.in.\*
- 4: gb.on.\*
- 5: gb.ov.\*
- 6: gb.pat.\*
- 7: gb.ph.\*
- 8: gb.pl.\*
- 9: gb.pr.\*
- 10: gb.ro.\*
- 11: gb.sts.\*
- 12: gb.sy.\*
- 13: gb.un.\*
- 14: gb.vi.\*
- 15: em.ba.\*
- 16: em.fun.\*
- 17: em.hum.\*
- 18: em.in.\*
- 19: em.mu.\*
- 20: em.om.\*
- 21: em.or.\*
- 22: em.ov.\*
- 23: em.pat.\*
- 24: em.ph.\*
- 25: em.pl.\*
- 26: em.ro.\*
- 27: em.sts.\*
- 28: em.un.\*
- 29: em.vi.\*
- 30: em.htg.hum.\*
- 31: em.htg.inv.\*
- 32: em.htg.other.\*
- 33: em.htg.mus.\*
- 34: em.htg.pln.\*
- 35: em.htg.rod.\*
- 36: em.htg.man.\*
- 37: em.htg.vrt.\*
- 38: em.sy.\*
- 39: em.htgo.hum.\*
- 40: em.htgo.mus.\*
- 41: em.htgo.other.\*

Pred. No. is the number of results predicted by chance to have a

score greater than or equal to the score of the result being printed,  
and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match %	Length	DB	ID	Description
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4	499.4	99.7	3769	9	AK096204	AK096204 Homo sapi
c 5	145.4	29.0	207387	2	AC115766	AC115766 Mus muscu
c 6	130.6	28.1	275631	2	AC125563	AC125563 Rattus no
7	113	22.6	618	9	AF493928	AF493928 Homo sapi
8	113	22.6	618	9	BT007025	BT007025 Homo sapi
9	113	22.6	618	12	BT007756	BT007756 Synthetic
10	113	22.6	800	6	AR270528	AR270528 Sequence
11	113	22.6	800	9	HSU27768	U27768 Human KGP4
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13	113	22.6	2753	9	BC051869	BC051869 Homo sapi
14	113	22.6	2934	6	AX451335	AX451335 Sequence
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16	100.2	20.0	1489	10	RNU27767	U27767 Rattus norv
17	100.2	20.0	2919	10	AF117211	AF117211 Rattus no
18	98.6	19.7	2781	10	BC003882	BC003882 Mus muscu
19	97	19.4	630	10	AB004315	AB004315 Mus muscu
20	89.8	17.9	201	10	RNU32327	U32327 Rattus norv
21	78.4	15.6	150	5	AF090091	AF090091 Gallus ga
22	76.8	15.3	254	5	AF090081	AF090081 Gallus ga
23	65	13.0	964	5	AF263451	AF263451 Xenopus l
24	54.2	10.8	275631	2	AC125563	AC125563 Rattus no
25	53.4	10.7	3653	9	AF009356	AF009356 Homo sapi
c 26	53.4	10.7	102812	2	AL158215	AL158215 Homo sapi
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30	52.6	10.5	192394	2	EX119907	EX119907 Danio rer
31	52.2	10.4	2078	4	SSC549925	AJ549925 Sus scrof
c 32	51.8	10.3	147047	9	AL353778	AL353778 Human DNA
33	51.6	10.3	6905	10	AY138504	AY138504 Mus muscu
c 34	51.6	10.3	50095	2	AC074333	AC074333 Mus muscu
c 35	51.6	10.3	132762	10	AL844529	AL844529 Mouse DNA
c 36	50.6	10.1	144798	2	AC141948	AC141948 Rattus no
c 37	50.6	10.1	188016	2	AC121256	AC121256 Mus muscu
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39	49.8	9.9	201	5	AB038435	AB038435 Xenopus l
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41	49	9.8	378	6	BD168779	BD168779 New disea
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ALIGNMENTS

RESULT 1	AX451337	Sequence 3 from Patent WO0216653.	20300 bp	DNA	linear	PAT 03-JUL-2002
LOCUS	AX451337	Sequence 3 from Patent WO0216653.				
DEFINITION	AX451337	Sequence 3 from Patent WO0216653.				
ACCESSION	AX451337	Sequence 3 from Patent WO0216653.				
VERSION	AX451337.1	GI:21698388				
KEYWORDS		synthetic construct				
SOURCE		synthetic construct				
ORGANISM		artificial sequences.				
REFERENCE	1	Levitt,P.R., Mirnics,K., Kodavali,V.C. and Nimgaonkar,V.L.				
AUTHORS		Methods and systems for facilitating the diagnosis and treatment of				
TITLE		schizophrenia				
JOURNAL		Patent: WO 0216653-A 3 28-FEB-2002;				





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QY 421 ACAAGCCGACATCTAGAGCTCAATACCTCTTGTGATGAGGCCAGAGAGATT 480
Db 87101 ACAAGCCGACATCTAGAGCTCAATACCTCTTGTGATGAGGCCAGAGAGATT 87160

QY 481 TTCAACCTGATGAGAGAGAT 501
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RESULT 3
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LOCUS
DEFINITION
Homo sapiens chromosome 1 clone RP11-288018, WORKING DRAFT
SEQUENCE, 3 unordered pieces.
AC031977
AC031977.7 GI:13194952
HTG: HTGS_PHASE1; HTGS_DRAFT; HTGS_FULLTOP; HTGS_ACTIVEFIN.
KEYWORDS
SOURCE
Homo sapiens
ORGANISM
Homo sapiens
REFERENCE
AUTHORS
Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 191699)
Abola, A.P., Bruno, D., Conn, L., Della Rosa, M., Faulkner, D.,
Fedorov, N., Glukhov, S., Hansen, N., Herman, Z.S., Hyman, R.,
Mao, J., Marathe, R., Morehouse, A.J., Oefner, P., Palm, C.J.,
Ramirez, D., Wilhelmy, J., Yu, S. and Davis, R.W.
Southwick, A.M., Webb, C., Wilhelmy, J., Yu, S. and Davis, R.W.
Unpublished
2 (bases 1 to 191699)
Abola, A.P., Bruno, D., Conn, L., Della Rosa, M., Faulkner, D.,
Fedorov, N., Glukhov, S., Hansen, N., Herman, Z.S., Hyman, R.,
Mao, J., Marathe, R., Morehouse, A.J., Oefner, P., Palm, C.J.,
Ramirez, D., Wilhelmy, J., Yu, S. and Davis, R.W.
Direct Submission
Submitted (03-APR-2000) DNA Sequencing and Technology Center,
Stanford University, 855 California Avenue, Palo Alto, CA 94304,
USA
On Mar 4, 2001 this sequence version replaced gi:9665085.
----- Genome Center
Center: Stanford DNA Sequencing and Technology Development
Center
Center code: SDSTDC
Web site: http://sequence-www.stanford.edu/group/human/
Contact: hum-info@sequence.stanford.edu
----- Project Information
Center project name: 880
Center clone name: RP11-288018
----- Summary Statistics
Sequencing Vector: M13mp18; X02513; 100% of reads
Sequencing Vector: plasmid; plasmid_accession; 0% of reads
Chemistry: Dye-primer; 1% of reads
Assembly program: Phrap; version 0.990319
Consensus quality: 190680 bases at least Q40
Consensus quality: 191287 bases at least Q30
Consensus quality: 191336 bases at least Q20
Insert size: 195548; agarose-fp
Insert size: 194499; sum-of-coverage
Quality coverage: 7.9x in Q20 bases; agarose-fp
Quality coverage: 8.1x in Q20 bases; sum-of-coverage.
NOTE: This is a 'working draft' sequence. It currently
consists of 3 contigs. The true order of the pieces
is not known and their order in this sequence record is
arbitrary. Gaps between the contigs are represented as
runs of N, but the exact sizes of the gaps are unknown.
This record will be updated with the finished sequence
as soon as it is available and the accession number will
be preserved.
1 12646: contig of 12646 bp in length
* 12647 12746: gap of unknown length
* 12747 94961: contig of 82215 bp in length
* 94962 95061: gap of unknown length

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FEATURES             * 95062 191699: contig of 96638 bp in length.
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Best Local Similarity 100.0%; Pred. No. 1.5e-138;
Matches 501; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
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Db 189481 TGGCAGAGAACTCTCTGGAATCTTAGTGAAGCTTCCTAGAAATAGTGGAGCTGACTATCATA 189422
QY 61 ATCTTGACAAACCCCAATAAATCAGATTTTAAATAATCTTTTATCCATGTGGCTTAC 120
Db 189421 ATCTTGACAAACCCCAATAAATCAGATTTTAAATAATCTTTTATCCATGTGGCTTAC 189362
QY 121 CATAACTCCCTGCATGAATTTTCTGATGAATCTCCCAATTTGTTTGAAGACAGAA 180
Db 189361 CATAACTCCCTGCATGAATTTTCTGATGAATCTCCCAATTTGTTTGAAGACAGAA 189302
QY 181 GATCTGCGCTGCTCTCTCTTAAAGCAGAAAGGTTTCATTCTGAACTTTTCTACTCTCTCA 240
Db 189301 GATCTGCGCTGCTCTCTCTTAAAGCAGAAAGGTTTCATTCTGAACTTTTCTACTCTCTCA 189242
QY 241 CATGTGCAAGGAGGACCCCAATGTCTACTTTGTTGTTTGTCTCTGAAATACAGAGGGTG 300
Db 189241 CATGTGCAAGGAGGACCCCAATGTCTACTTTGTTGTTTGTCTCTGAAATACAGAGGGTG 189182
QY 301 CACTGCACTTACAAGTCACTCAAGCCTACAGCTTGCATCTCTCAACGGGATATAGG 360
Db 189181 CACTGCACTTACAAGTCACTCAAGCCTACAGCTTGCATCTCTCAACGGGATATAGG 189122
QY 361 TCTAATGAAGCTTGGGCTTTGCCCTCAGGTGAACCTGGAATTTTGCACCCAGGGAAGAG 420
Db 189121 TCTAATGAAGCTTGGGCTTTGCCCTCAGGTGAACCTGGAATTTTGCACCCAGGGAAGAG 189062
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Db 189061 ACAAGCCGACATCTAGAGCTCAATACCTCTGCTTTGATGAGGCCAGAGAGATT 189002
QY 481 TTCAACCTGATGAGAGAGAT 501
Db 189001 TTCAACCTGATGAGAGAGAT 188981

RESULT 4
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LOCUS
DEFINITION
Homo sapiens cDNA FLJ38885 fis, clone MESAN2017417, moderately
similar to REGULATOR OF G-PROTEIN SIGNALING 4.
AC096204
AC096204.1 GI:21755635
VERSION
KEYWORDS
oligo capping; fis (full insert sequence).
SOURCE
Homo sapiens
ORGANISM
Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

```

REFERENCE  
AUTHORS  
1 Tanigami,A., Fujiwara,T., Shibahara,T., Goto,Y., Hirao,M., Shimizu,F., Wakebe,H., Ono,T., Hishigaki,H., Watanabe,T., Ozaki,K., Sugiyama,T., Irie,R., Otsuki,T., Sato,H., Wakamatsu,A., Ishii,S., Yamamoto,J., Isono,Y., Kawai-Hio,Y., Saito,K., Nishikawa,T., Kimura,K., Yamashita,H., Matsuo,K., Nakamura,Y., Sekine,M., Kikuchi,H., Kanda,K., Wagatsuma,M., Murakawa,K., Kanehori,K., Takahashi-Fujii,A., Oshima,A., Sugiyama,A., Kawakami,B., Suzuki,Y., Sugano,S., Nagahara,Y., Masuho,Y., Nagai,K. and Isogai,T.  
NEDO human cDNA sequencing project  
Unpublished  
2 (bases 1 to 3769)  
Isogai,T. and Yamamoto,J.  
Direct Submission  
TITLE  
JOURNAL  
AUTHORS  
TITLE  
JOURNAL  
COMMENT  
Kazusa-Kamatari, Kisarazu, Chiba 292-0812, Japan  
(E-mail: genomics@kri.co.jp, Tel:81-438-52-3975, Fax:81-438-52-3986)  
NEDO human cDNA sequencing project supported by Ministry of Economy, Trade and Industry of Japan; cDNA full insert sequencing: Research Association for Biotechnology (RAB); cDNA library construction: Helix Research Institute (HRI) (supported by Japan Key Technology Center etc.); 5'- & 3'-end one pass sequencing: RAB, HRI, and Biotechnology Center, National Institute of Technology and Evaluation; clone selection for full insert sequencing: HRI and RAB; annotation: HRI and RAB.  
Location/Qualifiers  
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Best Local Similarity 99.8%; Pred. No. 3.7e-138;  
Matches 500; Conservative 0; Mismatches 1; Indels 0; Gaps 0;  
QY 1 TGGCAGAGACTCTCGATCTTGTAGTGAAGTTCCTAGAGTGGAGCTGACTATCATTA 60  
DB 1114 TGGCAGAGACTCTCGATCTTGTAGTGAAGTTCCTAGAGTGGAGCTGACTATCATTA 1173  
QY 61 ATCTTGACACCCCAATAATCACTAGTTTAAAAAATCTCTTTATCCATGTGGCTTAC 120  
DB 1174 ATCTTGACACCCCAATAATCACTAGTTTAAAAAATCTCTTTATCCATGTGGCTTAC 1233  
QY 121 CATTAACCTCCCTGATGAATTTTCTGATGAATCTCCCAATTTGTTAGACAGAACGAA 180  
DB 1234 CATTAACCTCCCTGATGAATTTTCTGATGAATCTCCCAATTTGTTAGACAGAACGAA 1293  
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DB 1294 GATCTTGCCCTGCTCTCTTAAGCAGAAAGGTTTCATTCTGAACCTTTTCACTCTCTCA 1353  
QY 241 CATGTGCCAAGGAGGCCCAATCTCACTTTTGTGTTTGTCTTCTGAAATACAGAGGGTG 300  
DB 1354 CATGTGCCAAGGAGGCCCAATCTCACTTTTGTGTTTGTCTTCTGAAATACAGAGGGTG 1413  
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DB 1414 CACTTGCCACTTACAGTCACTACAAAGATACAGCTTGATCTCTCAACAGGGATATAGG 1473  
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DB 1534 ACAAGCCGGAACTGCTAGAGCCTACATTAACCTGCTTTGTATGAGGCCCAAGAGAGATT 1593

QY 481 TTCAACCTGATGGAGAGGAT 501  
DB 1594 TTCAACCTGATGGAGAGGAT 1614  
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Mus musculus clone RP23-82124, WORKING DRAFT SEQUENCE, 9 unordered pieces.  
AC115766  
AC115766 GI:29164623  
HTG; HTGS PHASE1; HTGS DRAFT.  
SOURCE  
Mus musculus (house mouse)  
ORGANISM  
Mus musculus  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.  
REFERENCE  
1 (bases 1 to 207387)  
Birren,B., Nusbaum,C. and Lander,E.  
Mus musculus, clone RP23-82124  
Unpublished  
2 (bases 1 to 207387)  
Birren,B., Linton,L., Nusbaum,C., Lander,E., Ali,A., Allen,N., Anderson,S., Barna,N., Bastien,V., Bloom,T., Boguslavsky,L., Boukhgalter,B., Brown,A., Camarata,J., Campopiano,A., Chang,J., Chazaro,B., Choepel,Y., Colangelo,M., Collins,S., Collymore,A., Cook,A., Cooke,P., DeArellano,K., Dewar,K., Diaz,J.S., Dodge,S., Faro,S., Ferreira,P., FitzHugh,W., Gage,D., Galagan,J., Gardyna,S., Ginde,S., Gord,S., Goyette,M., Graham,L., Grand-Pierre,N., Jones,C., Hagos,B., Horton,L., Hulme,W., Iliev,I., Johnson,R., Jones,C., Kamaat,A., Karatas,A., Kells,C., LaRocque,K., Lamazares,R., Lander,T., Lechoczky,J., Levine,R., Lindblad-Toh,K., Liu,G., MacLean,C., Macdonald,P., Major,J., Marquis,N., Matthews,C., McCarthy,M., McEwan,P., McKernan,K., Meldrim,J., Meneus,L., Mihova,T., Mlenga,V., Murphy,T., Naylor,J., Nguyen,C., Nicol,R., Norbu,C., Norman,C.H., O'Connor,T., O'Donnell,P., O'Neill,D., Oliver,J., Peterson,K., Phunkhang,P., Pierre,N., Pollara,V., Raymond,C., Retta,R., Rieback,M., Riley,R., Rise,C., Rogov,P., Roman,J., Rosetti,M., Roy,A., Santos,R., Schauer,S., Schuback,R., Seaman,S., Severy,P., Spencer,B., Stange-Thomann,N., Stojanovic,N., Straus,N., Subramanian,A., Talamas,J., Tesfaye,S., Theodore,J., Topham,K., Travers,M., Travis,N., Trigilio,J., Vassiliev,H., Viel,R., Vo,A., Wilson,B., Wu,X., Wyman,D., Ye,W.J., Young,G., Zainoun,J., Zembek,L., Zimmer,A. and Zody,M.  
Direct Submission  
TITLE  
JOURNAL  
REFERENCE  
AUTHORS  
Submitted (22-MAR-2002) Whitehead Institute/MIT Center for Genome Research, 320 Charles Street, Cambridge, MA 02141, USA  
3 (bases 1 to 207387)  
Birren,B., Nusbaum,C., Lander,E., Abouelleil,A., Allen,N., Anderson,S., Arachchi,H.M., Barna,N., Bastien,V., Bloom,T., Boguslavsky,L., Boukhgalter,B., Camarata,J., Chang,J., Choepel,Y., Collymore,A., Cook,A., Cooke,P., Corum,B., DeArellano,K., Diaz,J.S., Dodge,S., Dooley,K., Dorris,L., Erickson,J., Faro,S., Ferreira,P., FitzGerald,M., Gage,D., Galagan,J., Gardyna,S., Graham,L., Grand-Pierre,N., Hafez,N., Hagopian,D., Hagos,B., Hall,J., Horton,L., Hulme,W., Iliev,I., Johnson,R., Jones,C., Kamaat,A., Karatas,A., Kells,C., Lander,T., Levine,R., Lindblad-Toh,K., Liu,G., Lui,A., Mabbitt,R., MacLean,C., Macdonald,P., Major,J., Manning,J., Matthews,C., McCarthy,M., Meldrim,C., Nicol,R., Norbu,C., O'Connor,T., O'Donnell,P., O'Neill,D., Oliver,J., Peterson,K., Phunkhang,P., Pierre,N., Rachupka,A., Ramasamy,U., Raymond,C., Retta,R., Rise,C., Rogov,P., Roman,J., Schauer,S., Schuback,R., Seaman,S., Severy,P., Smith,C., Spencer,B., Stange-Thomann,N., Stojanovic,N., Stubbs,M., Talamas,J., Tesfaye,S., Theodore,J., Topham,K., Travers,M., Vassiliev,H., Venkataraman,V.S., Viel,R., Vo,A., Wilson,B., Wu,X., Wyman,D., Young,G., Zainoun,J., Zembek,L., Zimmer,A. and Zody,M.  
Direct Submission  
TITLE  
JOURNAL  
COMMENT  
Submitted (23-MAR-2003) Whitehead Institute/MIT Center for Genome Research, 320 Charles Street, Cambridge, MA 02141, USA  
On Mar 23, 2003 this sequence version replaced gi:28191504.

All repeats were identified using RepeatMasker:  
Smit, A.F.A. & Green, P. (1996-1997)  
<http://ftp.genome.washington.edu/RM/RepeatMasker.html>

----- Genome Center  
Center: Whitehead Institute/ MIT Center for Genome Research  
Center code: WIBR

Web site: <http://www-seq.wi.mit.edu>  
Contact: [sequence.submissions@genome.wi.mit.edu](mailto:sequence.submissions@genome.wi.mit.edu)

----- Project Information

Center project name: L2317

Center clone name: 82-I-24

----- Summary Statistics

Sequencing vector: Plasmid; n/a; 100% of reads

Chemistry: Dye-terminator Big Dye; 100% of reads

Assembly program: Phrap; version 0.960731

Consensus quality: 205497 bases at least Q40

Consensus quality: 206243 bases at least Q30

Consensus quality: 206432 bases at least Q20

Insert size: 205000; agarose-fp

Insert size: 206587; sum-of-contigs

Quality coverage: 8.8 in Q20 bases; agarose-fp

Quality coverage: 8.7 in Q20 bases; sum-of-contigs

\* NOTE: This is a 'working draft' sequence. It currently  
\* consists of 9 contigs. The true order of the pieces  
\* is not known and their order in this sequence record is  
\* arbitrary. Gaps between the contigs are represented as  
\* runs of N, but the exact sizes of the gaps are unknown.  
\* This record will be updated with the finished sequence  
\* as soon as it is available and the accession number will  
\* be preserved.

\* 1  
\* 61925: contig of 61925 bp in length  
\* 61926: gap of 100 bp  
\* 62026: contig of 2578 bp in length  
\* 64603: gap of 100 bp  
\* 64604: contig of 3873 bp in length  
\* 64703: gap of 100 bp  
\* 68576: gap of 100 bp  
\* 68577: contig of 3260 bp in length  
\* 71936: gap of 100 bp  
\* 71937: contig of 100 bp  
\* 72036: gap of 100 bp  
\* 72037: contig of 4533 bp in length  
\* 76569: gap of 100 bp  
\* 76570: contig of 5894 bp in length  
\* 82563: gap of 100 bp  
\* 82564: contig of 36987 bp in length  
\* 82664: gap of 100 bp  
\* 119651: contig of 100 bp  
\* 119751: contig of 45424 bp in length  
\* 165175: gap of 100 bp  
\* 165275: contig of 42113 bp in length.

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Best Local Similarity 74.3%; Pred. No. 3.7e-32;

Matches 197; Conservative 0; Mismatches 66; Indels 2; Gaps 1;

QY 239 CACATGTCGCAAGAGGAGGACCCCATGTCACATTTGTTTGTCTTCTGAAA--TACAGAG 296

Db 111101 CACATATGTGAGGCGACACCAATATGCCGGGTTTCATTCGGGGGGAATTAAGAA 111042

QY 297 GGTGCATCTGCACCTTACAGTCTACATAAGCATACAGGCTTGTCATCTCTCAACAGGGATA 356

Db 111041 AGGTCCAGCCACTCTCTGTTTCATTCAGGCTGAGCCCTCTCTTCAACAGGGCAC 110982

QY 357 TAGTCTTAATGAAGCCCTTGCCCTTCAGTGAAGTCTGATCTTGTGACACGGGA 416

Db 110981 TAGATCTAATGCAACCTTGGTCTTTGCACTTCAGGTGAACCTGCACTCTTGCACACAGA 110922

QY 417 AGAGCAGCCGGAACATGCTAGAGCTTACATAACCTGTTTGTATGAGGCCCAAGAGAA 476

Db 110921 GGAGCAGCCGGAACATGTTACAGCCACATAAATCTGTTTGTATGAGGCCCAAGAGAA 110862

QY 477 GATTTCACACCTGATGAGAGGAT 501

Db 110861 GATTTCACACCTGATGAGAGGAT 110837

## RESULT 6

## AC125563/c

## LOCUS

AC125563 275631 bp DNA linear HTG 09-NOV-2002  
Rattus norvegicus clone CH230-9B12, WORKING DRAFT SEQUENCE, 4

## DEFINITION

unordered pieces.

## ACCESSION

AC125563.4 GI:24817906

## VERSION

HTG; HTGS\_PHASE1; HTGS\_DRAFT; HTGS\_FULLTOP.

## KEYWORDS

Rattus norvegicus (Norway rat)

## SOURCE

Rattus norvegicus

## ORGANISM

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae;

Rattus.

1 (bases 1 to 275631)

## REFERENCE

## AUTHORS

Muzny, D. Marie, Metzker, M. Lee, Abramson, S., Adams, C., Alder, J.,

Allen, C., Allien, H., Alsbrooks, S., Amin, A., Anguiano, D.,

Anyalebechi, V., Ayvagi, A., Ayodeji, M., Baca, E., Baden, H.,

Baldwin, D., Bandaranaike, D., Barber, M., Barnstead, M., Benahmed, F.,

Biswal, K., Blair, J., Blankenburg, K., Blyth, P., Brown, M.,

Bryant, N., Buhay, C., Burch, P., Burrell, K., Calderon, E.,

Cardenas, V., Carter, K., Cavazos, I., Cesar, H., Center, A.,

Chacko, J., Chavez, D., Chen, G., Chen, R., Chen, Y., Chen, Z., Chu, J.,

Cleveland, C., Cockrell, R., Cox, C., Coyle, M., Cree, A., D'Souza, L.,

Davila, M. L., Davis, C., Davy-Carroll, L., De Anda, C., Dederich, D.,

Delgado, O., Denison, S., Deramo, C., Ding, Y., Dinh, H., Divya, K.,

Draper, H., Dugan-Rocha, S., Dunn, A., Durbin, K., Duval, B., Eaves, K.,

Egan, A., Escotto, M., Eugene, C., Evans, C. A., Falls, T., Fan, G.,

Fernandez, S., Finley, M., Flagg, N., Forbes, L., Foster, M., Foster, P.,

Fraser, C. M., Gabisi, A., Ganta, R., Garcia, A., Garner, T., Garner, W.,

Gebrgeorgis, E., Geer, K., Gill, R., Grady, M., Guerra, W., Guevara, W.,

Gunaratne, P., Haaland, W., Hamil, C., Hamilton, C., Hamilton, K.,

Harvey, Y., Havlak, P., Hawes, A., Henderson, N., Hernandez, J.,

Hernandez, R., Hines, S., Hladun, S. I., Hodgson, A., Hognes, M.,

Hollins, B., Howells, S., Hulyk, S., Hume, J., Idlebirt, D., Jackson, A.,

Jackson, L., Jacob, L., Jiang, H., Johnson, B., Johnson, R., Jolivet, A.,

Karpachy, S., Kelly, S., Khan, Z., King, L., Kovar, C.,

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Liu, J., Liu, W., Liu, Y., London, P., Longacre, S., Lopez, J.,

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Maheshwari, M., Mahdian, M., Mahmoud, M., Malloy, K., Mangum, A.,

Mangum, B., Mapua, P., Martin, K., Martin, R., Martinez, E.,

Mawhiney, S., McLeod, M.P., McNeill, T.Z., Meenen, E., Milosavljevic, A., Miner, G., Minja, E., Montemayor, J., Moore, S., Morgan, M., Morris, K., Morris, S., Munidasa, M., Murphy, M., Nair, L., Narkervis, C., Neal, D., Newton, N., Nguyen, N., Norris, S., Nwakoleneh, O., Okwuonu, G., Olarnpunsagoon, A., Pal, S., Parks, K., Pasternak, S., Paul, H., Perez, A., Perez, L., Pfannkuch, C., Plopper, F., Poldinger, A., Popovic, D., Primus, E., Pu, L.-L., Puzo, M., Quiróz, J., Rachlin, E., Reeves, K., Regier, M.A., Reigh, R., Reilly, B., Reilly, M., Ren, Y., Reuter, M., Richards, S., Riggs, F., Rives, C., Rodkey, T., Rojas, A., Rose, M., Rose, R., Ruiz, S.J., Sanders, M., Savary, G., Scherer, S., Scott, G., Shatsman, S., Shen, H., Shetty, J., Shvartsbeyn, A., Sisson, I., Sitter, C.D., Smajls, D., Sneed, A., Sodergren, E., Song, X.-Z., Sorelle, R., Sosa, J., Steimle, M., Strong, R., Sutton, A., Svatek, A., Taber, P., Taylor, C., Taylor, T., Thomas, N., Thomas, S., Tingey, A., Trejos, Z., Usmani, K., Vallas, R., Vera, V., Villasana, D., Waldron, L., Walker, B., Wang, J., Wang, Q., Wang, S., Warren, J., Warren, R., Wei, X., White, F., Williams, G., Willson, R., Wleczyk, R., Wooden, H., Worley, K., Wright, D., Wright, R., Wu, J., Yakub, S., Yen, J., Yoon, L., Yoon, V., Yu, F., Zhang, J., Zhou, J., Zhou, X., Zhao, S., Dunn, D., von Niederhausern, A., Weiss, R., Smith, D.R., Hoit, R.A., Smith, H.O., Weinstock, G. and Gibbs, R.A.

# TITLE JOURNAL

## REFERENCE AUTHORS

### TITLE JOURNAL

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Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 389 AGGTGAACCTGGATTCTTGACACGAGGAGAGACAGCGGACATGCTAGAGCCTACAA 448
Db 377 AGGTGAACCTGGATTCTTGACACGAGGAGAGACAGCGGACATGCTAGAGCCTACAA 436
QY 449 TAACCTGCTTTGATGAGGCCCAAGAGAGATTTCACCTGATGAGAGAGGAT 501
Db 437 TAACCTGCTTTGATGAGGCCCAAGAGAGATTTCACCTGATGAGAGAGGAT 489

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DEFINITION Homo sapiens regulator of G-protein signalling 4 mRNA, complete cds.
ACCESSION  BT007025.1  GI:30582888
VERSION     BT007025.1  GI:30582888
KEYWORDS    FLI CDNA.
SOURCE      Homo sapiens (human)
ORGANISM    Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 618)
Kainine,N., Chen,X., Rolfs,A., Halleck,A., Hines,L., Eisenstein,S.,
Koundinya,M., Raphael,J., Moreira,D., Kelley,T., LaBaer,J., Lin,Y.,
Phelan,M. and Farmer,A.
Cloning of human full-length CDSs in BD Creator(TM) System Donor
vector
Unpublished
2 (bases 1 to 618)
Kainine,N., Chen,X., Rolfs,A., Halleck,A., Hines,L., Eisenstein,S.,
Koundinya,M., Raphael,J., Moreira,D., Kelley,T., LaBaer,J., Lin,Y.,
Phelan,M. and Farmer,A.
Direct Submission
Submitted (13-MAY-2003) BD Biosciences Clontech, 1020 East Meadow
Circle, Palo Alto, CA 94303, USA
This CDS clone is a part of a collection of human full length
expression clones generated by BD Biosciences Clontech and the
Harvard Institute of Proteomics. Each CDS has been cloned in two
forms: with and without stop-codon (to allow fusion with C-terminal
tag). The CDS has been directionally cloned using BD In-Fusion(TM)
cloning system between the Sali and HindIII sites of the pDNR-DUAL
vector. Additional sequences in the clone: 'ACC' after Sali site
and before 'ATG' to provide kozak consensus sequence; 'GG' after
last codon and before HindIII site to maintain reading frame.
Clone distribution: http://bioinfo.clontech.com/orfclones.
Location/Qualifiers
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BASE COUNT      190 a 133 c 151 g 144 t
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Query Match      22.6%; Score 113; DB 9; Length 618;
Best Local Similarity 100.0%; Pred. No. 1.3e-22;
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QY 389 AGGTGAACCTGGATTCTTGACACGAGGAGAGACAGCGGACATGCTAGAGCCTACAA 448
Db 377 AGGTGAACCTGGATTCTTGACACGAGGAGAGACAGCGGACATGCTAGAGCCTACAA 436
QY 449 TAACCTGCTTTGATGAGGCCCAAGAGAGATTTCACCTGATGAGAGAGGAT 501
Db 437 TAACCTGCTTTGATGAGGCCCAAGAGAGATTTCACCTGATGAGAGAGGAT 489

RESULT 9
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DEFINITION Synthetic construct Homo sapiens regulator of G-protein signalling
4 mRNA, partial cds.
ACCESSION  BT007756.1  GI:30584350
VERSION     BT007756.1  GI:30584350
KEYWORDS    FLI CDNA.
SOURCE      synthetic construct
ORGANISM    synthetic construct
artificial sequences.
1 (bases 1 to 618)
Kainine,N., Chen,X., Rolfs,A., Halleck,A., Hines,L., Eisenstein,S.,
Koundinya,M., Raphael,J., Moreira,D., Kelley,T., LaBaer,J., Lin,Y.,
Phelan,M. and Farmer,A.
Cloning of human full-length CDSs in BD Creator(TM) System Donor
vector
Unpublished
2 (bases 1 to 618)
Kainine,N., Chen,X., Rolfs,A., Halleck,A., Hines,L., Eisenstein,S.,
Koundinya,M., Raphael,J., Moreira,D., Kelley,T., LaBaer,J., Lin,Y.,
Phelan,M. and Farmer,A.
Direct Submission
Submitted (13-MAY-2003) BD Biosciences Clontech, 1020 East Meadow
Circle, Palo Alto, CA 94303, USA
This CDS clone is a part of a collection of human full length
expression clones generated by BD Biosciences Clontech and the
Harvard Institute of Proteomics. Each CDS has been cloned in two
forms: with and without stop-codon (to allow fusion with C-terminal
tag). The CDS has been directionally cloned using BD In-Fusion(TM)
cloning system between the Sali and HindIII sites of the pDNR-DUAL
vector. Additional sequences in the clone: 'ACC' after Sali site
and before 'ATG' to provide kozak consensus sequence; 'GG' after
last codon and before HindIII site to maintain reading frame.
Clone distribution: http://bioinfo.clontech.com/orfclones.
Location/Qualifiers
1. .618
/organism="synthetic construct"
/mol_type="mRNA"
/db_xref="taxon:32630"
/clone="GH00329L1.0"
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SPKLSPKAKKIYNEFISVQATKEVNDLSDCTRETSRNMLEPTTCFDEAOKKIFNLM
EKDSYRFLKSRFYLDLVPSSCGAEKQKGAKSADCASLVPQCA"
BASE COUNT 189 a 133 c 151 g 145 t
ORIGIN
Query Match 22.6%; Score 113; DB 12; Length 618;
Best Local Similarity 100.0%; Pred. No. 1.3e-22;
Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 389 AGGTGAACCTGGATTCTTGACCCAGGGAAGAGACAGCCGGAACATGCTAGAGCTACAA 448
Db 377 AGGTGAACCTGGATTCTTGACCCAGGGAAGAGACAGCCGGAACATGCTAGAGCTACAA 436
Qy 449 TAACTGCTTTGATGAGGCCCGAGAGAGATTTTCAACTGATGAGAGAGAT 501
Db 437 TAACTGCTTTGATGAGGCCCGAGAGAGATTTTCAACTGATGAGAGAGAT 489

RESULT 10
AR270528 800 bp DNA linear PAT 10-APR-2003
LOCUS
DEFINITION Sequence 1091 from patent US 6500938.
ACCESSION AR270528
VERSION AR270528.1 GI:29701762
KEYWORDS
SOURCE
ORGANISM
Unknown.
Unclassified.
REFERENCE
1 (bases 1 to 800)
AUTHORS Au-Young,J. and Seilhamer,J.J.
TITLE Composition for the detection of signaling pathway gene expression
JOURNAL Patent: US 6500938-A 1091 31-DEC-2002;
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Location/Qualifiers
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Best Local Similarity 100.0%; Pred. No. 1.4e-22;
Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 389 AGGTGAACCTGGATTCTTGACCCAGGGAAGAGACAGCCGGAACATGCTAGAGCTACAA 448
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Qy 449 TAACTGCTTTGATGAGGCCCGAGAGAGATTTTCAACTGATGAGAGAGAT 501
Db 534 TAACTGCTTTGATGAGGCCCGAGAGAGATTTTCAACTGATGAGAGAGAT 586

RESULT 11
HSU27768 800 bp mRNA linear PRI 07-MAR-1996
LOCUS
DEFINITION Human RGP4 mRNA, complete cds.
ACCESSION U27768
VERSION U27768.1 GI:1216372
KEYWORDS
SOURCE
ORGANISM
Homo sapiens (human)
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
1 (bases 1 to 800)

```

```

AUTHORS Druey,K.M., Blumer,K.J., Kang,V.H. and Kehrl,J.H.
TITLE Inhibition of G-protein-mediated MAP kinase activation by a new
mammalian gene family
JOURNAL Nature 379 (6567), 742-746 (1996)
MEDLINE 96178495
PUBMED 8602223
REFERENCE 2 (bases 1 to 800)
AUTHORS Druey,K.
TITLE Direct Submission
JOURNAL Submitted (25-MAY-1995) Kirk Druey, Intramural Research/NIAD/LIR,
Rm 11B13, National Institutes of Health, 10 Center Drive, MSC 1876,
Bethesda, MD 20892-1876, USA
FEATURES
Location/Qualifiers
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/mol_type="mRNA"
/db_xref="taxon:9606"
/tissue_type="brain"
98..715
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/product="RGP4"
/protein_id="AAC50395.1"
/db_xref="GI:1216373"
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VVICQVSEVKKWASLENIISHEGLAFAKFLKSEYSEENIDFWISCEYKKIK
SPKLSPKAKKIYNEFISVQATKEVNDLSDCTRETSRNMLEPTTCFDEAOKKIFNLM
EKDSYRFLKSRFYLDLVPSSCGAEKQKGAKSADCASLVPQCA"
BASE COUNT 241 a 181 c 195 g 183 t
ORIGIN
Query Match 22.6%; Score 113; DB 9; Length 800;
Best Local Similarity 100.0%; Pred. No. 1.4e-22;
Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 389 AGGTGAACCTGGATTCTTGACCCAGGGAAGAGACAGCCGGAACATGCTAGAGCTACAA 448
Db 474 AGGTGAACCTGGATTCTTGACCCAGGGAAGAGACAGCCGGAACATGCTAGAGCTACAA 533
Qy 449 TAACTGCTTTGATGAGGCCCGAGAGAGATTTTCAACTGATGAGAGAGAT 501
Db 534 TAACTGCTTTGATGAGGCCCGAGAGAGATTTTCAACTGATGAGAGAGAT 586

RESULT 12
BC000737 1869 bp mRNA linear PRI 12-JUL-2001
LOCUS
DEFINITION Homo sapiens, regulator of G-protein signalling 4, clone MGC:2124
IMAGE:3510260, mRNA, complete cds.
ACCESSION BC000737
VERSION BC000737.1 GI:12653888
KEYWORDS
SOURCE
ORGANISM
Homo sapiens (human)
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
1 (bases 1 to 1869)
AUTHORS Strausberg,R.
TITLE Direct Submission
JOURNAL Submitted (15-NOV-2000) National Institutes of Health, Mammalian
Gene Collection (MGC), Cancer Genomics Office, National Cancer
Institute, 31 Center Drive, Room 11A03, Bethesda, MD 20892-2590,
USA
REMARK
COMMENT
NIH-MGC Project URL: http://mgc.nci.nih.gov
Contact: MGC help desk
Email: cgaps-r@mail.nih.gov
Tissue Procurement: ATCC
cDNA Library Preparation: Rubin Laboratory
cDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)
DNA Sequencing by: Institute for Systems Biology
http://www.systemsbio.org
contact: amadan@systemsbiology.org
Anup Madan, Rachel Dickhoff, Jessica Fahey, Stephanie Ford, Julia
Greene, Mark Kettman and Anuradha Madan

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Clone distribution: MGC clone distribution information can be found through the I.M.A.G.E. Consortium/LLNL at: <http://image.llnl.gov>  
Series: IRAL Plate: 7 Row: d Column: 22  
This clone was selected for full length sequencing because it passed the following selection criteria: matched mRNA gi: 1216372.

FEATURES

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/db\_xref="taxon:9606"  
/clone="MGC:2124 IMAGE:3510260"  
/tissue\_type="Brain, neuroblastoma"  
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/lab\_host="DH10B-R"  
/notes="Vector: pOTB7"  
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/protein\_id="AAH00737.1"  
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531 a 392 c 418 g 528 t

BASE COUNT

ORIGIN  
Query Match 22.6%; Score 113; DB 9; Length 1869;  
Best Local Similarity 100.0%; Pred. No. 1.4e-22;  
Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 389 AGTGAACTGGATTCTTCACCGAGGAGACGACCGGACATGCTAGACCTACAA 448  
Db |||||  
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Db |||||  
QY 449 TAACCTGCTTTGATGAGGCCAGAGAAGATTTCACCTGATGGAGAGGAT 501  
Db |||||  
QY 522 TAACCTGCTTTGATGAGGCCAGAGAAGATTTCACCTGATGGAGAGGAT 574  
Db |||||

RESULT 13

BC051869  
LOCUS  
DEFINITION Homo sapiens regulator of G-protein signalling 4, mRNA (CDNA clone  
MGC:60244 IMAGE:5786695), complete cds.  
ACCESSION BC051869  
VERSION BC051869.1 GI:30354011  
KEYWORDS MGC.  
SOURCE Homo sapiens (human)  
ORGANISM Homo sapiens

REFERENCE

1 (bases 1 to 2753)  
Strausberg, R.L., Feingold, E.A., Grouse, L.H., Derge, J.G., Klausner, R.D., Collins, F.S., Wagner, L., Shenmen, C.M., Schuler, G.D., Altschul, S.F., Zeeberg, B., Buetow, K.H., Schaefer, C.F., Bhat, N.K., Hopkins, R.F., Jordan, H., Moore, T., Max, S.I., Wang, J., Hsieh, F., Dietzenko, L., Marusina, K., Farmer, A.A., Rubin, G.M., Hong, L., Stapleton, M., Soares, M.B., Bonaldo, M.F., Casavant, T.L., Scheetz, T.E., Brownstein, M.J., Umed, T.B., Toshiyuki, S., Carninci, P., Prange, C., Raha, S., Loquellano, N.A., Peters, G.J., Abramson, R.D., Mullany, S.J., Bosak, S.A., McSwan, P.J., McKernan, K.J., Malek, J.A., Gunaratne, P.H., Richards, S., Worley, K.C., Hale, S., Garcia, A.M., Gay, L.J., Hulyk, S.W., Villalón, D.K., Muzny, D.M., Sodergren, E.J., Lu, X., Gibbs, R.A., Sahney, J., Helton, E., Kettman, M., Madan, A., Rodriguez, S., Sanchez, A., Whitting, M., Madan, A., Young, A.C., Shevchenko, Y., Bouffard, G.G., Blakesley, R.W., Touchman, J.W., Green, E.D., Dickson, M.C., Rodriguez, A.C., Grimwood, J., Schmutz, J., Myers, R.M., Butterfield, Y.S., Krzywinski, M.I., Skalska, U., Small, D.E., Schnerch, A., Schein, J.E., Jones, S.J. and Marra, M.A.

TITLE  
JOURNAL  
MEDLINE  
PUBMED  
REFERENCE  
AUTHORS  
TITLE  
JOURNAL  
REMARK  
COMMENT  
Generation and initial analysis of more than 15,000 full-length human and mouse cDNA sequences  
Proc. Natl. Acad. Sci. U.S.A. 99 (26), 16899-16903 (2002)  
22388257  
12477932  
2 (bases 1 to 2753)  
Strausberg, R.  
Direct Submission  
Submitted (01-MAY-2003) National Institutes of Health, Mammalian Gene Collection (MGC), Cancer Genomics Office, National Cancer Institute, 31 Center Drive, Room 11A03, Bethesda, MD 20892-2590, USA  
NIH-MGC Project URL: <http://mgc.nci.nih.gov>  
Contact: MGC help desk  
Email: [cgaps-remail.nih.gov](mailto:cgaps-remail.nih.gov)  
Tissue Procurement: ATCC  
cDNA Library Preparation: Life Technologies, Inc.  
cDNA Library Arrayed by: The I.M.A.G.E. Consortium (LLNL)  
DNA Sequencing by: Sequencing Group at the Stanford Human Genome Center, Stanford University School of Medicine, Stanford, CA 94305  
Web site: <http://www-shgc.stanford.edu>  
Contact: (Dickson, Mark) [mcd@paxil.stanford.edu](mailto:mcd@paxil.stanford.edu)  
Dickson, M., Schmutz, J., Grimwood, J., Rodriguez, A., and Myers, R. M.

Clone distribution: MGC clone distribution information can be found through the I.M.A.G.E. Consortium/LLNL at: <http://image.llnl.gov>  
Series: IRAL Plate: 110 Row: g Column: 22  
This clone was selected for full length sequencing because it passed the following selection criteria: matched mRNA gi: 11184227.

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784 a 549 c 570 g 850 t

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QY 474 AGTGAACTGGATTCTTCACCGAGGAGACGACCGGACATGCTAGACCTACAA 533  
Db |||||  
QY 449 TAACCTGCTTTGATGAGGCCAGAGAAGATTTCACCTGATGGAGAGGAT 501  
Db |||||  
QY 534 TAACCTGCTTTGATGAGGCCAGAGAAGATTTCACCTGATGGAGAGGAT 586  
Db |||||

BASE COUNT

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Query Match 22.6%; Score 113; DB 9; Length 2753;  
Best Local Similarity 100.0%; Pred. No. 1.5e-22;  
Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 389 AGTGAACTGGATTCTTCACCGAGGAGACGACCGGACATGCTAGACCTACAA 448  
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QY 474 AGTGAACTGGATTCTTCACCGAGGAGACGACCGGACATGCTAGACCTACAA 533  
Db |||||  
QY 449 TAACCTGCTTTGATGAGGCCAGAGAAGATTTCACCTGATGGAGAGGAT 501  
Db |||||  
QY 534 TAACCTGCTTTGATGAGGCCAGAGAAGATTTCACCTGATGGAGAGGAT 586  
Db |||||  
RESULT 14  
AX451335  
LOCUS AX451335 2934 bp DNA linear PAT 03-JUL-2002

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DEFINITION Sequence 1 from Patent WO0216653.
ACCESSION AX451335
VERSION AX451335.1 GI:21698387
SOURCE Homo sapiens (human)
ORGANISM Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE 1
AUTHORS Levitt,P.R., Mirnics,K., Kodavali,V.C. and Nimgaonkar,V.L.
TITLE Methods and systems for facilitating the diagnosis and treatment of
schizophrenia
JOURNAL Patent: WO 0216653-A 1 28-FEB-2002;
UNIVERSITY University of Pittsburgh (US)
FEATURES
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Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 399 AGGTGAACCTGGATTCTTGACACAGGGAAGAGACAGCGGAAACATGCTAGAGCCTACAA 448
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QY 449 TAACTGCTTTGATGAGGCCAGAGAGAGATTTTCAACCTGATGAGAGGAT 501
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Db 513 TAACTGCTTTGATGAGGCCAGAGAGAGATTTTCAACCTGATGAGAGGAT 565
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RESULT 15
ACCESSION AK093959
LOCUS Homo sapiens cDNA FLJ36640 fis, clone TRACH2019151, moderately
similar to REGULATOR OF G-PROTEIN SIGNALING 4.
ACCESSION AK093959
VERSION AK093959.1 GI:21752924
KEYWORDS oligo capping; fis (full insert sequence).
SOURCE Homo sapiens (human)
ORGANISM Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE 1
AUTHORS Suzuki,O., Sasaki,N., Aotsuka,S., Shoji,T., Ichihara,T.,
Shiohata,N., Matsumoto,K., Hirano,M., Sano,S., Nomura,R.,
Yoshikawa,Y., Matsumura,Y., Moriya,S., Chiba,E., Momiyama,H.,
Onogawa,S., Kaeriyama,S., Satoh,N., Matsunawa,H., Takahashi,E.,
Katsoka,R., Kuga,N., Kuroda,A., Satoh,I., Kamata,K., Takami,S.,
Terashima,Y., Watanabe,M., Sugiyama,T., Irie,R., Otsuki,T.,
Sato,H., Ota,T., Wakamatsu,A., Ishii,S., Yamamoto,J., Isono,Y.,
Kawai-Hio,Y., Saito,K., Nishikawa,T., Kimura,K., Yamashita,H.,
Matsuo,K., Nakamura,Y., Sekine,M., Kikuchi,H., Kanda,K.,
Wagatsuma,M., Murakawa,K., Kanehori,K., Takahashi-Fujii,A.,
Oshima,A., Sugiyama,A., Kawakami,B., Suzuki,Y., Sugano,S.,
Nagahara,K., Masuho,Y., Nagai,K. and Isogai,T.
TITLE NEDO human cDNA sequencing project
JOURNAL Unpublished
REFERENCE 2 (bases 1 to 2599)
AUTHORS Isogai,T. and Yamamoto,J.
TITLE Direct Submission
JOURNAL Submitted (04-JUL-2002) Takao Isogai, FLJ Project (HRI Team); 2-6-7
Kazusa-Kamatari, Kisarazu, Chiba 292-0812, Japan
(E-mail:genomics@ri.co.jp, Tel:81-438-52-3975, Fax:81-438-52-3986)
COMMENT NEDO human cDNA sequencing project supported by Ministry of
Economy, Trade and Industry of Japan; cDNA full insert sequencing:
Research Association for Biotechnology (RAB); cDNA library
construction: Helix Research Institute (HRI) (supported by Japan
Key Technology Center etc.); 5'- & 3'-end one pass sequencing: RAB,

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HRI, and Biotechnology Center, National Institute of Technology and
Evaluation; clone selection for full insert sequencing: HRI and
RAB; annotation: HRI and RAB.
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BASE COUNT 719 a 520 c 541 g 819 t
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Best Local Similarity 94.3%; Pred. No. 3.3e-22;
Matches 116; Conservative 0; Mismatches 7; Indels 0; Gaps 0;

QY 379 TTGCCCCTCAGGTGAACCTGGATTCTTGACACAGGGAAGAGACAGCGGAAACATGCTA 438
|||||
Db 324 TTAGTCATGAATGTGAACCTGGATTCTTGACACAGGGAAGAGACAGCGGAAACATGCTA 383
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QY 439 GAGCCTACAATAACCTGCTTTGATGAGGCCAGAGAGAGATTTTCAACCTGATGGAGAAG 498
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Db 384 GAGCCTACAATAACCTGCTTTGATGAGGCCAGAGAGAGATTTTCAACCTGATGGAGAAG 443
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QY 499 GAT 501
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Db 444 GAT 446

Search completed: November 7, 2003, 11:11:24
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GenCore version 5.1.6  
Copyright (c) 1993 - 2003 CompuGen Ltd.

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Run on: November 7, 2003, 01:54:16 ; Search time 1962.4 Seconds  
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10444.222 Million cell updates/sec

Title: US-09-939-209A-3\_COPY\_19800\_20300  
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Sequence: 1 ccacatgattatctcaatag.....atgagtgaactccattccac 501

Scoring table: IDENTITY\_NUC  
Gapop 10.0 , Gapext 1.0

Searched: 2888711 seqs, 2045481386 residues

Total number of hits satisfying chosen parameters: 5777422

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

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2: gb\_htg:\*  
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32: em\_htg\_mus:\*  
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40: em\_htgo\_mus:\*  
41: em\_htgo\_other:\*

Pred. No. is the number of results predicted by chance to have a

score greater than or equal to the score of the result being printed,  
and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB	ID	Description
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3	496.2	99.0	191699	2	AC031977	AC031977 Homo sapi
4	481.8	96.2	116800	9	AL391379	AL391379 Human DNA
5	480.2	95.8	165295	2	AC068486	AC068486 Homo sapi
6	480.2	95.8	194867	2	AC023152	AC023152 Homo sapi
7	478.6	95.5	149428	2	AC010264	AC010264 Homo sapi
8	478.6	95.5	163314	2	AL590557	AL590557 Homo sapi
9	478.6	95.5	173158	2	AC090096	AC090096 Homo sapi
10	478.6	95.5	177097	2	AP001569	AP001569 Homo sapi
11	478.6	95.5	179726	9	AC007052	AC007052 Homo sapi
12	478.6	95.5	182411	2	AC090408	AC090408 Homo sapi
13	478.6	95.5	186780	9	AC005740	AC005740 Homo sapi
14	478.6	95.5	196869	2	AC087535	AC087535 Homo sapi
15	478.6	95.5	199275	2	AC011401	AC011401 Homo sapi
16	478.6	95.5	200774	2	AP001592	AP001592 Homo sapi
17	477	95.2	21399	9	AL310112	AL310112 Homo sapi
18	477	95.2	41389	9	AP000542	AP000542 Homo sapi
19	477	95.2	48103	9	AL356282	AL356282 Human DNA
20	477	95.2	60451	9	AL392112	AL392112 Human DNA
21	477	95.2	77127	9	AC104065	AC104065 Homo sapi
22	477	95.2	78735	9	AC008545	AC008545 Homo sapi
23	477	95.2	84579	2	AL356380	AL356380 Homo sapi
24	477	95.2	94882	2	AC002317	AC002317 Homo sapi
25	477	95.2	97352	9	AL441885	AL441885 Human DNA
26	477	95.2	97771	9	BX088563	BX088563 Human DNA
27	477	95.2	109140	2	AP001969	AP001969 Homo sapi
28	477	95.2	112084	9	AC104648	AC104648 Homo sapi
29	477	95.2	118879	9	AC106854	AC106854 Homo sapi
30	477	95.2	132337	9	AC104798	AC104798 Homo sapi
31	477	95.2	136198	9	AC092092	AC092092 Homo sapi
32	477	95.2	142080	9	AC079748	AC079748 Homo sapi
33	477	95.2	144136	2	AC012243	AC012243 Homo sapi
34	477	95.2	153624	9	CNS01D84	AL121654 BAC seque
35	477	95.2	158091	2	AC025255	AC025255 Homo sapi
36	477	95.2	159698	9	AC027269	AC027269 Homo sapi
37	477	95.2	162746	9	AC006050	AC006050 Homo sapi
38	477	95.2	165047	9	AC079835	AC079835 Homo sapi
39	477	95.2	172579	9	AC036125	AC036125 Homo sapi
40	477	95.2	172600	9	AL359644	AL359644 Human DNA
41	477	95.2	173728	9	AL365496	AL365496 Human DNA
42	477	95.2	176597	2	AC080190	AC080190 Homo sapi
43	477	95.2	177176	2	AC040995	AC040995 Homo sapi
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ALIGNMENTS

RESULT 1  
AX451337  
LOCUS  
DEFINITION  
ACCESSION  
VERSION  
KEYWORDS  
SOURCE  
ORGANISM  
REFERENCE  
AUTHORS  
TITLE  
JOURNAL

AX451337  
Sequence 3 from Patent WO0216653.  
AX451337  
AX451337.1 GI:21698388  
synthetic construct  
synthetic construct  
artificial sequences.  
1  
Levitt,P.R., Mirmics,K., Kodavali,V.C. and Ningaonkar,V.L.  
Methods and systems for facilitating the diagnosis and treatment of  
schizophrenia  
Patent: WO 0216653-A 3 28-FEB-2002;

20300 bp  
DNA  
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PAT 03-JUL-2002

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Db 19800 CCACATGATTATCTCAATAGATGCGAGAAAGGCAATTTGACAAAATTTTAAACACTCTTCAT 19859
Qy 61 GCTAAAACCTCTCAATCAATAGTATTGATGGGACGTATCTCAAAAATAATAAGCACTAT 120
Db 19860 GCTAAAACCTCTCAATCAATAGTATTGATGGGACGTATCTCAAAAATAATAAGCACTAT 19919
Qy 121 CTATGACAACTCACAGCAATATCATCTGAATGGGCAAAAACCTGGAGCATTCCTCTT 180
Db 19920 CTATGACAACTCACAGCAATATCATCTGAATGGGCAAAAACCTGGAGCATTCCTCTT 19979
Qy 181 GAAACGGGCAAGACAGAGGATGCCCTCTCTCACCACTCTCTATTCACATAGTGTGGA 240
Db 19980 GAAACGGGCAAGACAGAGGATGCCCTCTCTCACCACTCTCTATTCACATAGTGTGGA 20039
Qy 241 AGCTCTGCCAGGGCAATAGGAGGAGAGAAATTAAGGGTATTCATTTAGGAGAGA 300
Db 20040 AGCTCTGCCAGGGCAATAGGAGGAGAGAAATTAAGGGTATTCATTTAGGAGAGA 20099
Qy 301 GGAAGTCAAAATGTCCTGTTTGCAGATGACATGATCTATATCTAGAAAACCCCATCGT 360
Db 20100 GGAAGTCAAAATGTCCTGTTTGCAGATGACATGATCTATCTAGAAAACCCCATCGT 20159
Qy 361 CTCAGCCCAAAATCTCTTAAGCTGATGAAGCAACTTCAGCAAGTCTCAGGATACAAAT 420
Db 20160 CTCAGCCCAAAATCTCTTAAGCTGATGAAGCAACTTCAGCAAGTCTCAGGATACAAAT 20219
Qy 421 CAATGTACAAAATCACAGCACTTTATATCATCAATTAACAGACAGAGGCCAAAT 480
Db 20220 CAATGTACAAAATCACAGCACTTTATATCATCAATTAACAGACAGAGGCCAAAT 20279
Qy 481 CATGAGTGAACTCCCATTCAC 501
Db 20280 CATGAGTGAACTCCCATTCAC 20300

RESULT 2
AL583850
LOCUS      Human DNA sequence from clone Rp11-430G6 on chromosome 1, complete
DEFINITION
ACCESSION  AL583850
VERSION     AL583850.5 GI:16973044
KEYWORDS   HTG.
SOURCE     Homo sapiens (human)
ORGANISM   Homo sapiens
            Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
            Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
            Tracey, A.
            Direct Submission
            Submitted (15-NOV-2001) Wellcome Trust Sanger Institute, Hinxton,
            Cambridgeshire, CB10 1SA, UK. E-mail enquiries:
            humquies@sanger.ac.uk
            On Nov 16, 2001 this sequence version replaced gi:15020514.
            During sequence assembly data is compared from overlapping clones.
```

```
Where differences are found these are annotated as variations
together with a note of the overlapping clone name. Note that the
variation annotation may not be found in the sequence submission
corresponding to the overlapping clone, as we submit sequences with
only a small overlap as described above.
This sequence was finished as follows unless otherwise noted: all
regions were either double-stranded or sequenced with an alternate
chemistry or covered by high quality data (i.e., phred quality >=
30); an attempt was made to resolve all sequencing problems, such
as compressions and repeats; all regions were covered by at least
one plasmid subclone or more than one M13 subclone; and the
assembly was confirmed by restriction digest. The following
abbreviations are used to associate primary accession numbers given
in the feature table with their source databases: Em, EMBL; Sw,
SWISSPROT; Tr, TREMBL; Wp, WORMPEP; Information on the WORMPEP
database can be found at
http://www.sanger.ac.uk/Projects/C_elegans/wormpep
This sequence
was generated from part of bacterial clone contigs of human
chromosome 1, constructed by the Sanger Centre Chromosome 1 Mapping
Group. Further information can be found at
http://www.sanger.ac.uk/HGP/Chrl
Rp11-430G6 is from the library RPCI-11.2 constructed by the group
of Pieter de Jong. For further details see
http://www.chori.org/bacpac/home.htm
VECTOR: pBACe3.6
IMPORTANT: This sequence is not the entire insert of clone
Rp11-430G6. It may be shorter because we sequence overlapping
sections only once, except for a short overlap.
The true right end of clone Rp11-430G6 is at 165329 in this
sequence. The true right end of clone Rp11-33H2 is at 2000 in this
sequence.
FEATURES
  source
    1. .165329
    /organism="Homo sapiens"
    /mol_type="genomic DNA"
    /db_xref="taxon:9606"
    /chromosome="1"
    /clone="Rp11-430G6"
    /clone_lib="RPCI-11.2"
BASE COUNT    51144 a 30897 c 31439 g 51849 t
ORIGIN
    Query Match      100.0%; Score 501; DB 9; Length 165329;
    Best Local Similarity 100.0%; Pred. No. 5.8e-114;
    Matches 501; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qy 1 CCACATGATTATCTCAATAGATGCGAGAAAGGCAATTTGACAAAATTTTAAACACTCTTCAT 60
Db 91481 CCACATGATTATCTCAATAGATGCGAGAAAGGCAATTTGACAAAATTTTAAACACTCTTCAT 91540
Qy 61 GCTAAAACCTCTCAATCAATAGTATTGATGGGACGTATCTCAAAAATAATAAGCACTAT 120
Db 91541 GCTAAAACCTCTCAATCAATAGTATTGATGGGACGTATCTCAAAAATAATAAGCACTAT 91600
Qy 121 CTATGACAACTCACAGCAATATCATCTGAATGGGCAAAAACCTGGAGCATTCCTCTT 180
Db 91601 CTATGACAACTCACAGCAATATCATCTGAATGGGCAAAAACCTGGAGCATTCCTCTT 91660
Qy 181 GAAACGGGCAAGACAGAGGATGCCCTCTCTCACCACTCTCTATTCAACATAGTGTGGA 240
Db 91661 GAAACGGGCAAGACAGAGGATGCCCTCTCTCACCACTCTCTATTCAACATAGTGTGGA 91720
Qy 241 AGCTCTGGCAGGGCAATTAGGAGGAGAGAAATTAAGGGTATTCATTTAGGAGAGA 300
Db 91721 AGCTCTGGCAGGGCAATTAGGAGGAGAGAAATTAAGGGTATTCATTTAGGAGAGA 91780
Qy 301 GGAAGTCAAAATGTCCTGTTTGCAGATGACATGATTTGTATATCTAGAAAACCCCATCGT 360
Db 91781 GGAAGTCAAAATGTCCTGTTTGCAGATGACATGATTTGTATATCTAGAAAACCCCATCGT 91840
Qy 361 CTCAGCCCAAAATCTCTTAAGCTGATGAAGCAACTTCAGCAAGTCTCAGGATACAAAT 420
Db 91841 CTCAGCCCAAAATCTCTTAAGCTGATGAAGCAACTTCAGCAAGTCTCAGGATACAAAT 91900
```

QY 421 CCAATGTACAAAATCAAGCACTCTTATACATCAATAACAGCAAAACAGAGGCAAAAT 480  
 Db 91901 CCAATGTACAAAATCAAGCACTCTTATACATCAATAACAGCAAAACAGAGGCAAAAT 91960  
 QY 481 CATGAGTGAACCTCCCATTCAC 501  
 Db 91961 CATGAGTGAACCTCCCATTCAC 91981

RESULT 3  
 AC031977/c  
 LOCUS  
 DEFINITION Homo sapiens chromosome 1 clone RP11-288018, WORKING DRAFT  
 SEQUENCE, 3 unordered pieces.  
 AC031977 GI:13194952  
 HTG; HTGS\_PHASE1; HTGS\_DRAFT; HTGS\_FULLTOP; HTGS\_ACTIVEFIN.  
 SOURCE Homo sapiens (human)  
 ORGANISM Homo sapiens  
 REFERENCE  
 AUTHORS Abola, A.P., Bruno, D., Conn, L., Della Rosa, M., Faulkner, D.,  
 Federspiel, N., Glukhov, S., Hansen, N., Herman, Z.S., Hyman, R.,  
 Mao, J., Komp, C., Kottler, S., Lam, B., Marathe, R., Miranda, M.,  
 Morehouse, A.J., Nguyen, M., Oefner, P., Palm, C.J., Ramirez, D.,  
 Southwick, A.M., Webb, C., Wilhelmy, J., Yu, S. and Davis, R.W.  
 Unpublished  
 JOURNAL  
 REFERENCE 2 (bases 1 to 191699)  
 AUTHORS Abola, A.P., Bruno, D., Conn, L., Della Rosa, M., Faulkner, D.,  
 Federspiel, N., Glukhov, S., Hansen, N., Herman, Z.S., Hyman, R.,  
 Mao, J., Marathe, R., Morehouse, A.J., Oefner, P., Palm, C.J.,  
 Ramirez, D., Wilhelmy, J., Yu, S. and Davis, R.W.  
 Direct Submission  
 TITLE Submitted (03-APR-2000) DNA Sequencing and Technology Center,  
 JOURNAL Stanford University, 855 California Avenue, Palo Alto, CA 94304,  
 USA  
 COMMENT On Mar 4, 2001 this sequence version replaced gi:9665085.

----- Genome Center  
 Center: Stanford DNA Sequencing and Technology Development  
 Center  
 Center code: SDS/DC  
 Web site: <http://sequence-www.stanford.edu/group/human/>  
 Contact: [hum-info@sequence.stanford.edu](mailto:hum-info@sequence.stanford.edu)  
 ----- Project Information  
 Center project name: 880  
 Center clone name: RP11-288018  
 ----- Summary Statistics

Sequencing Vector: M13mp18; X02513; 100% of reads  
 Sequencing Vector: plasmid; plasmid\_accession; 0% of reads  
 Chemistry: Dye-primer; 1% of reads  
 Chemistry: Dye-terminator Big Dye; 99% of reads  
 Assembly program: Phrap; version 0.990319  
 Consensus quality: 190660 bases at least Q40  
 Consensus quality: 191287 bases at least Q30  
 Consensus quality: 191336 bases at least Q20  
 Insert size: 195548; agarose-fp  
 Insert size: 191499; sum-of-ctgigs  
 Quality coverage: 7.9x in Q20 bases; agarose-fp  
 Quality coverage: 8.1x in Q20 bases; sum-of-ctgigs.

\* NOTE: This is a 'working draft' sequence. It currently  
 \* consists of 3 contigs. The true order of the pieces  
 \* is not known and their order in this sequence record is  
 \* arbitrary. Gaps between the contigs are represented as  
 \* runs of N, but the exact sizes of the gaps are unknown.  
 \* This record will be updated with the finished sequence.  
 \* as soon as it is available and the accession number will  
 \* be preserved.

\* 1 12646: contig of 12646 bp in length  
 \* 12647 12746: gap of unknown length  
 \* 12747 94961: contig of 82215 bp in length  
 \* 94962 95061: gap of unknown length

FEATURES \* 95062 191699: contig of 96638 bp in length.

Location/Qualifiers  
 source 1..191699  
 /organism="Homo sapiens"  
 /mol\_type="genomic DNA"  
 /db\_xref="taxon:9606"  
 /chromosome="1"  
 /clone\_lib="RPC1 human BAC library 11"  
 /clone 1..12646  
 misc\_feature  
 /note="assembly\_name:Contig5"  
 misc\_feature  
 /note="assembly\_name:Contig6"  
 clone\_end:17"  
 misc\_feature  
 /note="assembly\_name:Contig7"  
 /note="end:SP6"  
 clone\_end:SP6"

BASE COUNT 57654 a 36385 c 36166 g 61293 t 201 others  
 ORIGIN

Query Match 99.0%; Score 496.2; DB 2; Length 191699;  
 Best Local Similarity 99.4%; Pred. No. 8.8e-113;  
 Matches 498; Conservative 0; Mismatches 3; Indels 0; Gaps 0;  
 QY 1 CCACATGATTATCTCAATAGATGCAGAAAGCGCATTTGACAAAATTTTAAACAACCTCTTCAT 60  
 Db 184686 CCACATGATTATCTCAATAGATGCAGAAAGCGCTTTGACAAAATTTTAAACAACCTCTTCAT 184627  
 QY 61 GCTAAAACTCTCAATCAATTTAGTATTTGATGGACGCTATCTCAAAAATAATAAGCACTAT 120  
 Db 184626 GCTAAAACTCTCAATCAATTTAGTATTTGATGGACGCTATCTCAAAAATAATAAGCACTAT 184567  
 QY 121 CTATGACAACTCAGCCCAATATCATCTCAATGGGCAAAAACCTGGAGCAATTCCTCTTT 180  
 Db 184566 CTATGACAACTCAGCCCAATATCATCTCAATGGGCAAAAACCTGGAGCAATTCCTCTTT 184507  
 QY 181 GAAAACGGGCGACAGACAGGATGCCCTCTCTCAACCACTCTCTCAACCTCTCTCAACATAGTGTGGA 240  
 Db 184506 GAAAACGGGCGACAGACAGGATGCCCTCTCTCAACCACTCTCTCAACCTCTCTCAACATAGTGTGGA 184447  
 QY 241 AGCTGTGCCCGAGGCAATTTAGGCGAGAGAAATAAAGGGTATTTCAATTAGGAGAGA 300  
 Db 184446 AGCTGTGCCCGAGGCAATTTAGGCGAGAGAAATAAAGGGTATTTCAATTAGGAGAGA 184387  
 QY 301 GGAAGTCAAAATGTCCCTGTTTGCAGATGACATGATGTTATCTAGAAAACCCCATCGT 360  
 Db 184386 GGAAGTCAAAATGTCCCTGTTTGCAGATGACATGATGTTATCTAGAAAACCCCATCGT 184327  
 QY 361 CTCAGCCCAAAATCTCTTAAAGCTGATAAGCAACTTCAGCAAGTCTCAGGATCAAAAT 420  
 Db 184326 CTCAGCCCAAAATCTCTTAAAGCTGATAAGCAACTTCAGCAAGTCTCAGGATCAAAAT 184267  
 QY 421 CAATGTACAAAATCAACAGCACTCTTATCATCAATAACAGCAAAACAGAGGCCAAAT 480  
 Db 184266 CAATGTACAAAATCAACAGCACTCTTATCATCAATAACAGCAAAACAGAGGCCAAAT 184207  
 QY 481 CATGAGTGAACCTCCCATTCAC 501  
 Db 184206 CATGAGTGAACCTCCCATTCAC 184186

RESULT 4  
 AL391379/c

LOCUS Human DNA sequence from clone RP13-171J5 on chromosome X, complete  
 DEFINITION sequence.

ACCESSION AL391379

VERSION AL391379.12 GI:13560021

KEYWORDS HTG.

SOURCE Homo sapiens (human)

ORGANISM

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
 Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

## REFERENCE

1. (bases 1 to 116800)  
 Chapman, J.  
 Direct Submission  
 Submitted (05-APR-2001) Sanger Centre, Hinxton, Cambridgeshire,  
 CB10 1SA, UK. E-mail enquiries: humquery@sanger.ac.uk  
 requests: clonerequest@sanger.ac.uk  
 On Apr 6, 2001 this sequence version replaced gi:13446478.  
 During sequence assembly data is compared from overlapping clones.  
 Where differences are found these are annotated as variations  
 together with a note of the overlapping clone name. Note that the  
 variation annotation may not be found in the sequence submission  
 corresponding to the overlapping clone, as we submit sequences with  
 only a small overlap as described above.  
 This sequence was finished as follows unless otherwise noted: all  
 regions were either double-stranded or sequenced with an alternate  
 chemistry or covered by high quality data (i.e., phred quality >= 30); an attempt was made to resolve all sequencing problems, such  
 as compressions and repeats; all regions were covered by at least  
 one plasmid subclone or more than one M13 subclone; and the  
 assembly was confirmed by restriction digest. The following  
 abbreviations are used to associate primary accession numbers given  
 in the feature table with their source databases: Em, EMBL; Sw,  
 SWISSPROT; Tr, TrEMBL; Wp, WORMPEP; Information on the WORMPEP  
 database can be found at  
[http://www.sanger.ac.uk/Projects/C\\_elegans/wormpep](http://www.sanger.ac.uk/Projects/C_elegans/wormpep) This sequence  
 was generated from part of bacterial clone contigs of human  
 chromosome X, constructed by the Sanger Centre Chromosome X Mapping  
 Group. Further information can be found at  
<http://www.sanger.ac.uk/HGP/ChrX>  
 RP13-171J5 is from the library RPCI-13.1 constructed by the group  
 of Pieter de Jong. For further details see  
<http://www.chori.org/bacpac/home.htm>  
 VECTOR: pBAC3.6

IMPORTANT: This sequence is not the entire insert of clone  
 RP13-171J5. It may be shorter because we sequence overlapping  
 sections only once, except for a 100 base overlap.  
 The true right end of clone RP13-171J5 is at 11680 in this  
 sequence. The true right end of clone RP11-348F1 is at 100 in this  
 sequence.

## FEATURES

## Source

## Location/Qualifiers

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1..116800
/organism="Homo sapiens"
/mol_type="genomic DNA"
/db_xref="taxon:9606"
/chromosome="X"
/clone="RP13-171J5"
/clone_lib="RPCI-13.1"
135..1785
/feature="L1P4 repeat: matches 4494..6146 of consensus"
1924..1979
/feature="L2 repeat: matches 2693..2748 of consensus"
2397..2867
/feature="L1MB1 repeat: matches 5693..6164 of consensus"
2888..2954
/feature="TH1B repeat: matches 328..394 of consensus"
2965..3777
/feature="L1A13 repeat: matches -651..189 of consensus"
3803..4094
/feature="L1P13 repeat: matches 288..606 of consensus"
4109..4774
/feature="L1P13 repeat: matches 758..1353 of consensus"
4775..5070
/feature="AluX repeat: matches 1..306 of consensus"
5071..5332
/feature="L1A13 repeat: matches 1353..1632 of consensus"
5348..8788
/feature="L1P repeat: matches 1794..5302 of consensus"
8789..8916
/feature="WSTB repeat: matches 1..112 of consensus"
8919..9466
/feature="L1MB repeat: matches 5090..5675 of consensus"
9528..9653
/feature="L1A1H repeat: matches 319..451 of consensus"

```

```

repeat_region
/feature="L1MA3 repeat: matches 6289..6304 of consensus"
9663..9679
/feature="L1MA3 repeat: matches 1..308 of consensus"
9680..9995
/feature="AluX repeat: matches 5527..6289 of consensus"
9996..10726
/feature="L1MA3 repeat: matches 4887..5298 of consensus"
10728..11133
/feature="L1ME repeat: matches 2929..3347 of consensus"
11130..11545
/feature="L1M1 repeat: matches 1..294 of consensus"
11546..11841
/feature="AluX repeat: matches 1029..2929 of consensus"
11842..13729
/feature="L1M1 repeat: matches 1380..1029 of consensus"
13730..14034
/feature="AluX repeat: matches 16965..17477 of consensus"
14035..16479
/feature="L1M1 repeat: matches 5401..6143 of consensus"
16965..17477
/feature="L1MA2 repeat: matches 53..166 of consensus"
17481..18227
/feature="L1MA2 repeat: matches 2179..2500 of consensus"
18228..20880
/feature="L2 repeat: matches 2617..2750 of consensus"
20881..21429
/feature="L2 repeat: matches 46..214 of consensus"
21430..21593
/feature="L1M1 repeat: matches 45..230 of consensus"
21594..23742
/feature="L1MA6 repeat: matches 4524..6143 of consensus"
23743..25363
/feature="L1P3 repeat: matches 6071..6146 of consensus"
25364..25803
/feature="L2 repeat: matches 2567..2746 of consensus"
25804..26189
/feature="L1M1 repeat: matches 188..262 of consensus"
26190..27355
/feature="L1MC3 repeat: matches 5355..7736 of consensus"
27356..29011
/feature="L1M4 repeat: matches 2942..3407 of consensus"
29012..30525
/feature="L1M4 repeat: matches 1..291 of consensus"
30526..30881
/feature="AluX repeat: matches 1864..2123 of consensus"
30882..31259
/feature="L1MEC repeat: matches 6155..6290 of consensus"
31260..32051
/feature="L1MA5 repeat: matches 1271..1908 of consensus"
32052..32293
/feature="L1M4C repeat: matches 979..1176 of consensus"
32294..33260
/feature="L1M4C repeat: matches 57..398 of consensus"
33261..34684
/feature="SVA repeat: matches 16..955 of consensus"
34685..35218
/feature="CpG island"
35219..35393
/feature="not experimental"
35394..35561
/feature="SVA repeat: matches 724..1386 of consensus"
35562..35561

```





```

* 13501 13600: gap of unknown length
* 13601 19166: contig of 5566 bp in length
* 19167 19266: gap of unknown length
* 19267 25083: contig of 5817 bp in length
* 25084 23183: gap of unknown length
* 25184 34033: contig of 8850 bp in length
* 34034 34133: gap of unknown length
* 34134 46088: contig of 11955 bp in length
* 46089 46188: gap of unknown length
* 46189 65638: contig of 19450 bp in length
* 65639 65738: gap of unknown length
* 65739 93577: contig of 27839 bp in length
* 93578 93677: gap of unknown length
* 93678 120965: contig of 27288 bp in length
* 120966 121065: gap of unknown length
* 121066 165295: contig of 44230 bp in length.
FEATURES
    source
        1..165295
            /organism="Homo sapiens"
            /mol_type="genomic DNA"
            /db_xref="taxon:9606"
            /chromosome="X"
            /clone="RP11-31605"
        1..1758
            /note="assembly_name:Contig3"
        1859..3835
            /note="assembly_name:Contig4"
            clone_end:T7
            vector_side:right
        3936..6268
            /note="assembly_name:Contig5"
        6369..9527
            /note="assembly_name:Contig6"
        9628..13500
            /note="assembly_name:Contig7"
        13601..19166
            /note="assembly_name:Contig8"
        19267..25083
            /note="assembly_name:Contig9"
            clone_end:SP6
            vector_side:left
        25184..34033
            /note="assembly_name:Contig10"
        34134..46088
            /note="assembly_name:Contig11"
        46189..65638
            /note="assembly_name:Contig12"
        65739..93577
            /note="assembly_name:Contig13"
        93678..120965
            /note="assembly_name:Contig14"
        121066..165295
            /note="assembly_name:Contig15"
BASE COUNT 50879 a 32297 c 32077 g 48840 t 1202 others
ORIGIN
Query Match          95.8%; Score 480.2; DB 2; Length 165295;
Best Local Similarity 97.4%; Pred. No. 8.1e-109;
Matches 488; Conservative 0; Mismatches 13; Indels 0; Gaps 0;
Qy 1 CCACATGATTATCTCAATAGATGCGAGAAAGGCATTTCACAAAATTTAAACACTCTTCAT 60
Db 83733 CCACATGATTATCTCAATAGATGCGAGAAAGGCCTTTCACAAAATTTAAACACGGTTTCA 83792
Qy 61 GCTAAAACTCTCAATCAATTAGGTATTGATGGGACGTATCTCAAAATAATAAGCACTAT 120
Db 83793 GCTAAAACTCTCAATAAATTAGGTATTGATGGGACGTATCTCAAAATAATAAGCACTAT 83852
Qy 121 CTATGACAACTCACAGCAATATCATCTAGTGGGCAAAAACCTGGGAAGCATTCCTTT 180
Db 83853 CTATGACAACTCACAGCAATATCATCTAGTGGGCAAAAACCTGGGAAGCATTCCTTT 83912
Qy 181 GAAACGGGCGACAGACAGGGATGCCCTCTCTCACCACTCTCTTCAACATAGTGTGGA 240

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---

```

Db 83913 GAAAACTGGCAAGACAGGGATGCCCTCTCTCACCACTCTCTTCAACATAGTGTGGA 83972
Qy 241 AGCTTCGCCAGGCAATTAGGCAGGAGGAAGGAATAAAGGGTATTCAATTAGGAGAAGA 300
Db 83973 AGTTTCGCCAGGCAATTAGGCAGGAGGAAGGAATAAAGGGTATTCAATTAGGAGAAGA 84032
Qy 301 GGAAGTCAAAATGTCCCTGTTTGAGATGACATGATTTGATATCTAGAAAACCCCATCGT 360
Db 84033 GGAAGTCAAAATGTCCCTGTTTGAGATGACATGATTTGATATCTAGAAAACCCCATGT 84092
Qy 361 CTCAGCCCAAAATCTCTTAAGCTGATAAGCACTTCAGCAAGTCTCAGATACAAAAT 420
Db 84093 CTCAGCCCAAAATCTCTTAAGCTGATAAGCACTTCAGCAAGTCTCAGATACAAAAT 84152
Qy 421 CAATGTACAAAATCAACAAGCACTTATATCATCAATAACAGACAGAGAGGCAAAAT 480
Db 84153 CAATGTACAAAATCAACAAGCACTTATATCATCAATAACAGACAGAGAGGCAAAAT 84212
Qy 481 CATGAGTGAATCCCATTCAC 501
Db 84213 CATGAGTGAATCCCATTCAC 84233

RESULT 6
AC023152 194867 bp DNA linear HTG 07-JUL-2000
LOCUS
DEFINITION Homo sapiens chromosome X clone RP11-737B10, WORKING DRAFT
SEQUENCE, 3 unordered pieces.
ACCESSION AC023152.3 GI:7191122
VERSION HTG; HTGS PHASE1; HTGS_DRAFT.
KEYWORDS Homo sapiens (human)
SOURCE Homo sapiens
ORGANISM Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE 1 (bases 1 to 194867)
AUTHORS Waterston,R.H.
TITLE The sequence of Homo sapiens clone
JOURNAL Unpublished
REFERENCE 2 (bases 1 to 194867)
AUTHORS Waterston,R.H.
TITLE Direct Submission
JOURNAL Submitted (08-FEB-2000) Genome Sequencing Center, Washington
University School of Medicine, 4444 Forest Park Parkway, St. Louis,
MO 63108, USA
COMMENT On Mar 7, 2000 this sequence version replaced gi:7140268.

----- Genome Center -----
Center: Washington University Genome Sequencing Center
Center code: WUGSC
Web site:http://genome.wustl.edu/gsc/index.shtml
----- Project Information -----
Center project name: H NH0737B10
----- Summary Statistics -----
Sequencing vector: M13; 100%
Sequencing vector: plasmid; 0%
Chemistry: Dye-primer ET; 100% of reads
Chemistry: Dye-terminator Big Dye; 0% of reads
Assembly program: Phrap; version 0.990319
Consensus quality: 192787 bases at least Q40
Consensus quality: 193180 bases at least Q30
Consensus quality: 193457 bases at least Q20
Insert size: 190000; agarose-fp
Insert size: 194667; sum-of-contigs
Quality coverage: 7.06 in Q20 bases; agarose-fp
Quality coverage: 6.92 in Q20 bases; sum-of-contigs
-----
* NOTE: This is a 'working draft' sequence. It currently
* consists of 3 contigs. The true order of the pieces
* is not known and their order in this sequence record is
* arbitrary. Gaps between the contigs are represented as
* runs of N, but the exact sizes of the gaps are unknown.

```

```
* This record will be updated with the finished sequence
* as soon as it is available and the accession number will
* be preserved.
*
* 1 8464: contig of 8464 bp in length
* 8465 8564: gap of unknown length
* 8565 78379: contig of 69815 bp in length
* 78380 78479: gap of unknown length
* 78480 194867: contig of 116388 bp in length.
FEATURES
    source
        1..194867
            /organism="Homo sapiens"
            /mol_type="genomic DNA"
            /db_xref="taxon:9606"
            /chromosome="X"
            /clone="RP11-737B10"
        1..8464
            /note="assembly_name:Contig4
            clone end:77
            vector side:right"
        8565..78379
            /note="assembly_name:Contig5
            clone end:SP6
            vector side:right"
        78480..194867
            /note="assembly_name:Contig6"
            /note="assembly_name:Contig6"
BASE COUNT 60197 a 38604 c 39004 g 56862 t 200 others
ORIGIN
    Query Match          95.8%; Score 480.2; DB 2; Length 194867;
    Best Local Similarity 97.4%; Pred. No. 8e-109;
    Matches 488; Conservative 0; Mismatches 13; Indels 0; Gaps 0;
QY 1 CCACATGATTATCTCAATAGATGCGAGAAAGGCATTGACAAAATTTAACACTTTCAT 60
Db 126543 CCACATGATTATCTCAATAGATGCGAGAAAGGCCTTGACAAAATTTCAACACGCTTCAT 126602
QY 61 GCTAAAACTCTCAATCAATAGGATTGATGGAGCGTATCTCAAAAATTAAGCACTAT 120
Db 126603 GCTAAAACTCTCAATCAATAGGATTGATGGAGCGTATCTCAAAAATTAAGCACTAT 126662
QY 121 CTATGACAACTCAGCAATATCATACTGAATGGGCAAAACTGGAAGCATTCCTCTT 180
Db 126663 CTATGACAACTCAGCAATATCATACTGAATGGGCAAAACTGGAAGCATTCCTCTT 126722
QY 181 GAAACGGGCAAGAGGATGCCCTCTCTACCACTCCTTATTCACATAGTGTGGA 240
Db 126723 GAAACGGGCAAGAGGATGCCCTCTCTACCACTCCTTATTCACATAGTGTGGA 126782
QY 241 AGCTCTGCCAGGCGCAATTAGGCGAGAGGAAGAAATAAGGTTATTCATTAAGGAGAGA 300
Db 126783 AGTTCTGCCAGGCGCAATTAGGCGAGAGGAAGAAATAAGGTTATTCATTAAGGAGAGA 126842
QY 301 GGAAGTCAAAATGTCCTCTTTGCGAGATGACATGTTGTATCTAGAAAAACCCCATCGT 360
Db 126843 GGAAGTCAAAATGTCCTCTTTGCGAGATGACATGTTGTATCTAGAAAAACCCCATGTT 126902
QY 361 CTCAGCCCAAAATCTCTTAGCTGATGAAGCACTTCAGCAAGTCTCAGATACAAAAT 420
Db 126903 CTCAGCCCAAAATCTCTTAGCTGATGAAGCACTTCAGCAAGTCTCAGATACAAAAT 126962
QY 421 CAATGTACAAAATCAACAAGCACTTATATCATCAATAACAGACAAAACAGAGGCCAAAT 480
Db 126963 CAATGTACAAAATCAACAAGCACTTATATCAACCAATTAACAGACAAAACAGAGGCCAAAT 127022
QY 481 CATGAGTGAATCCCATTCAC 501
Db 127023 CATGAGTGAATCCCATTCAC 127043
RESULT 7
AC010264/c 149428 bp DNA linear HTG 18-JUL-2000
LOCUS Homo sapiens chromosome 5 clone CTC-468K18, WORKING DRAFT SEQUENCE,
DEFINITION
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23 ordered pieces.
AC010264
AC010264.5 GI:9256174
HTG; HTGS PHASE2; HTGS_DRAFT.
SOURCE
Homo sapiens (human)
ORGANISM
Eukaryota; Metazoa; Chordata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
1 (bases 1 to 149428)
DOE Joint Genome Institute.
Sequencing of Human Chromosome 5
Unpublished
2 (bases 1 to 149428)
DOE Joint Genome Institute.
Direct Submission
Submitted (15-SEP-1999) Production Sequencing Facility, DOE Joint
Genome Institute, 2800 Mitchell Drive, Walnut Creek, CA 94598, USA
On Jul 18, 2000 this sequence version replaced gi:7710802.
-----Genome Center
Center: Joint Genome Institute
Center Code: JGI
Web site: http://www.jgi.doe.gov
-----
Project Information
Center Project Name: 365150
Center clone name: CIT-HSPC_468K18
-----
Summary Statistics
Consensus quality: 136337 bases at least Q40
Consensus quality: 144306 bases at least Q30
Consensus quality: 146065 bases at least Q20
Estimated insert size: 150000; pulse field gel estimation
Quality coverage: 148378; sum-of-contigs estimation
Quality coverage: 4.59 in Q20 bases; pulse field gel estimation
Quality coverage: 4.64 in Q20 bases; sum-of-contigs estimation.
* NOTE: This is a 'working draft' sequence. It currently
* consists of 23 contigs. Gaps between the contigs
* are represented as runs of N. The order of the pieces
* is believed to be correct as given, however the sizes
* of the gaps between them are based on estimates that have
* provided by the submitter.
* This sequence will be replaced
* by the finished sequence as soon as it is available and
* the accession number will be preserved.
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* 1 8196: contig of 8196 bp in length
* 8197 8296: gap of unknown length
* 8297 12916: contig of 4820 bp in length
* 12917 13016: gap of unknown length
* 13017 14756: contig of 1740 bp in length
* 14757 14856: gap of unknown length
* 14857 19143: contig of 4287 bp in length
* 19144 19243: gap of unknown length
* 19244 20796: contig of 1553 bp in length
* 20797 20897: gap of unknown length
* 20897 28283: contig of 7387 bp in length
* 28284 28384: gap of unknown length
* 28384 43450: contig of 15067 bp in length
* 43451 43551: gap of unknown length
* 43551 55287: contig of 11737 bp in length
* 55288 55388: gap of unknown length
* 55388 73601: contig of 18214 bp in length
* 73602 73701: gap of unknown length
* 73702 78295: contig of 4594 bp in length
* 78296 78395: gap of unknown length
* 78396 81832: contig of 3437 bp in length
* 81833 81932: gap of unknown length
* 81933 83998: contig of 2066 bp in length
* 83999 84099: gap of unknown length
* 84099 86531: contig of 2433 bp in length
* 86532 86632: gap of unknown length
* 86632 88757: contig of 2126 bp in length
* 88758 95081: contig of 6224 bp in length
* 88858
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* 95082 95181: gap of unknown length
* 95182 98288: contig of 3107 bp in length
* 98289 98388: gap of unknown length
* 98389 107715: contig of 9327 bp in length
* 107716 107815: gap of unknown length
* 107816 112037: contig of 4222 bp in length
* 112038 112137: gap of unknown length
* 112138 114033: contig of 1896 bp in length
* 114034 114133: gap of unknown length
* 114134 119274: contig of 5141 bp in length
* 119275 119374: gap of unknown length
* 119375 126879: contig of 7505 bp in length
* 126880 126979: gap of unknown length
* 126980 129104: contig of 2125 bp in length
* 129105 129205: gap of unknown length
* 129205 149428: contig of 20224 bp in length.

FEATURES             Location/Qualifiers
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                     /mol_type="genomic DNA"
                     /db_xref="taxon:9606"
                     /chromosome="5"
                     /clone="CTC-468K18"
                     /clone_lib="CalTech human BAC library C"
BASE COUNT    39562 a 32858 c 33987 g 40817 t 2204 others
ORIGIN
Query Match      95.5%; Score 478.6; DB 2; Length 149428;
Best Local Similarity 97.2%; Pred. No. 2e-108;
Matches 487; Conservative 0; Mismatches 14; Indels 0; Gaps 0;

Qy 1 CCACATGATTATCTCAATAGTAGTCAGAAAAGGCATTTACAAAATTTTAACAACTCTTCAT 60
Db 26772 CCACATGATTATCTCAATAGTAGTCAGAAAAGGCCTTTACAAAATTTCAACAACTTCAT 26713

Qy 61 GCTAAAAAATCTCAATCAATAGTATTGATGGGACGATATCTCAAAATTAATAAGCACTAT 120
Db 26712 GCTAAAAAATCTCAATCAATAGTATTGATGGGACGATATCTCAAAATTAATAAGCACTAT 26653

Qy 121 CTATGACAAATCTACGCAATATCATACTGAATGGGCAAAAACATGGAAGCATTCCTTTT 180
Db 26652 CTATGACAAATCTACGCAATATCATACTGAATGGGCAAAAACATGGAAGCATTCCTTTT 26593

Qy 181 GAAACGGGCACAGACAGGATGCCCTCTCTCACCCTCTATTCACATAGTGTGGA 240
Db 26592 GAAACCTGGCACAAGACAGGATGCCCTCTCTCACCCTCTATTCACATAGTGTGGA 26533

Qy 241 AGCTCTGCCAGGGCAATTAGGCAGGAGAGAAATTAAGGGTATTCATATTAGGAGAAGA 300
Db 26532 AGTTCTGCCAGGGCAATTAGGCAGGAGAGAAATTAAGGGTATTCATATTAGGAGAAGA 26473

Qy 301 GGAAGTCAAAATGTCCCTGTTTGCAGATGACATGATTTGTATATCTAGAAAAACCCCATCGT 360
Db 26472 GGAAGTCAAAATGTCCCTGTTTGCAGATGACATGATTTGTATATCTAGAAAAACCCCATGT 26413

Qy 361 CTCAGCCCAAAATCTCCCTTAAGCTGATAAGCAACTTCAGCAAGTCTCAGATACAAAT 420
Db 26412 CTCAGCCCAAAATCTCCCTTAAGCTGATAAGCAACTTCAGCAAGTCTCAGATACAAAT 26353

Qy 421 CAATGTACAAAATACAAAGCACTTTATACATCAATTAACAGACAAAACAGAGAGCCAAAT 480
Db 26352 CAATGTACAAAATACAAAGCACTTTATACATCAATTAACAGACAAAACAGAGAGCCAAAT 26293

Qy 481 CATGAGTCAACTCCCAATTCAC 501
Db 26292 CATGAGTCAACTCCCAATTCAC 26272

RESULT 8
LOCUS      AL590557/c      163314 bp      DNA      linear      HTG 20-JUL-2001
DEFINITION Homo sapiens chromosome 1 clone RP11-24C8, *** SEQUENCING IN
            PROGRESS ***, 8 unordered pieces.

```

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AL590557      AL590557.8      GI:13992136
VERSION      HTG; HTGS PHASE1; HTGS_CANCELLED.
KEYWORDS
SOURCE      Homo sapiens (human)
ORGANISM      Homo sapiens
              Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
              Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE      1
              Mclay, K.
              Direct Submission
              Submitted (20-JUL-2001) Sanger Centre, Hinxton, Cambridgeshire,
              CB10 1SA, UK. E-mail enquiries: humquery@sanger.ac.uk
              requests: clonerequest@sanger.ac.uk
              On May 8, 2001 this sequence version replaced gi:13990622.
              ----- Genome Center
              Center: Sanger Centre
              Center code: SC
              Web site: http://www.sanger.ac.uk
              Contact: humquery@sanger.ac.uk
              ----- Project Information
              Center project name: BA24C8
              ----- Summary Statistics
              Assembly program: XGAP4; version 4.5
              Sequencing vector: plasmid; L08752; 100% of reads
              Chemistry: Dye-terminator Big Dye; 100% of reads
              Consensus quality: 161385 bases at least Q40
              Consensus quality: 161940 bases at least Q30
              Consensus quality: 162264 bases at least Q20
              Insert size: 162614; sum-of-contigs
              Insert size: 164357; 4.9% error; agarose-fp
              Quality coverage: 6.63x in Q20 bases; sum-of-contigs Quality
              coverage: 6.67x in Q20 bases; agarose-fp
              -----
              * NOTE: This is a 'working draft' sequence. It currently
              * consists of 8 contigs. The true order of the pieces
              * is not known and their order in this sequence record is
              * arbitrary. Gaps between the contigs are represented as
              * runs of N, but the exact sizes of the gaps are unknown.
              * This record will be updated with the finished sequence
              * as soon as it is available and the accession number will
              * be preserved.
              *
              * 16091: contig of 16091 bp in length
              * 16092 16191: gap of 100 bp
              * 16192 51184: contig of 34993 bp in length
              * 51185 51284: gap of 100 bp
              * 51285 67941: contig of 16657 bp in length
              * 67942 68041: gap of 100 bp
              * 68042 103921: contig of 35880 bp in length
              * 103922 104021: gap of 100 bp
              * 104022 107020: contig of 2399 bp in length
              * 107021 107120: gap of 100 bp
              * 107121 123748: contig of 16628 bp in length
              * 123749 123848: gap of 100 bp
              * 123849 137091: contig of 13243 bp in length
              * 137092 137191: gap of 100 bp
              * 137192 163314: contig of 26123 bp in length.

FEATURES             Location/Qualifiers
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                     /organism="Homo sapiens"
                     /mol_type="genomic DNA"
                     /db_xref="taxon:9606"
                     /chromosome="1"
                     /clone="RP11-24C8"
                     /clone_lib="RPCI-11.1"
                     /note="assembly fragment:00603
                     fragment_chain:1"
   misc_feature       1..16091
                     /note="assembly fragment:01328
                     fragment_chain:1"
   misc_feature       16192..51184
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   misc_feature       51285..67941
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QY      421 CAATGTACAAAATCACAAGCACTCTTATACATCAATTAACAGACAAAACAGAGAGCCAAAT 480
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Db      29804 CAATGTACAAAATCACAAGCACTCTTATACACCAATAACAGACAAAACAGAGAGCCAAAT 29863
      |||||||
QY      481 CATGAGTGAATCCCAATTCAC 501
      |||||||
Db      29864 CATGAGTGAATCCCAATTCAC 29884
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RESULT 10
AP001569
LOCUS   AP001569 177097 bp DNA linear HTG 30-MAY-2000
DEFINITION Homo sapiens chromosome 18 clone RP11-859C21 map 18q21, WORKING
DRAFT SEQUENCE, 32 unordered pieces.
ACCESSION AP001569
VERSION   1
KEYWORDS  HTG; HTGS_PHASE1; HTGS_DRAFT.
SOURCE   Homo sapiens (human)
ORGANISM Homo sapiens

REFERENCE
1 (Bases 1 to 177097)
Hattori, M., Ishii, K., Toyoda, A., Taylor, T.D., Hong-Seog, P.,
Fujiyama, A., Yada, T., Totoki, Y., Watanabe, H. and Sakaki, Y.
Homo sapiens 177,097 genomic DNA of 18q21
Published Only in Database (2000)
2 (Bases 1 to 177097)
Hattori, M., Ishii, K., Toyoda, A., Taylor, T.D., Hong-Seog, P.,
Fujiyama, A., Yada, T., Totoki, Y., Watanabe, H. and Sakaki, Y.
Direct Submission
Submitted (29-MAR-2000) Masahira Hattori, The Institute of Physical
and Chemical Research (RIKEN), Genomic Sciences Center (GSC);
Kiyasato Univ., 1-15-1 Kitasato, Sagami-hara, Kanagawa 228-8555,
Japan (E-mail: hattori@gsc.riken.go.jp,
URL: http://hgp.gsc.riken.go.jp/, Tel: 81-42-778-9923,
Fax: 81-42-778-9924)
On May 30, 2000 this sequence version replaced gi:7380904.

----- Genome Center
Center: RIKEN Genomic Sciences Center (GSC)
Center code: RIKEN
Web site: http://hgp.gsc.riken.go.jp/
Contact: hattori@gsc.riken.go.jp
----- Project Information
Center project name: HumDrafl18
Center clone name: RP11-859C21
----- Summary Statistics
Sequencing vector: PCR products; 100% of reads
Chemistry: Dye-terminator ET-amersham; 100% of reads
Assembly program: Phrap; version 0.990329
Consensus quality: 160670 bases at least Q40
Consensus quality: 168058 bases at least Q30
Consensus quality: 171631 bases at least Q20
Insert size: 173997; sum-of-contigs
Quality coverage: 4.63x in Q20 bases; sum-of-contigs

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NOTE: This is a 'working draft' sequence. It currently consists of
32 contigs. The true order of the pieces is not known and their
order in this sequence record is arbitrary. Gaps between the
contigs are represented as runs N, but the exact sizes of the gaps
are unknown. This record will be updated with the finished sequence
as soon as it is available and the accession number will be
preserved
1 23792 contig of 23792 bp in length
23893 45465 contig of 21573 bp in length
45566 58167 contig of 12602 bp in length
58268 71173 contig of 9980 bp in length
91181 97387 contig of 6207 bp in length
97488 104902 contig of 7415 bp in length
105003 11107 contig of 6105 bp in length
111208 116851 contig of 5644 bp in length
116852 123585 contig of 4255 bp in length
123586 123685 contig of 6634 bp in length
123686 128985 contig of 5300 bp in length
128986 129085 contig of 100 bp in length
129086 134233 contig of 5148 bp in length
134234 134334 contig of 100 bp in length
134334 138588 contig of 4255 bp in length
138589 138688 contig of 100 bp in length
138689 142573 contig of 3885 bp in length
142574 142673 contig of 100 bp in length
142674 146286 contig of 3613 bp in length
146287 146386 contig of 100 bp in length
146387 150053 contig of 3667 bp in length
150054 150153 contig of 100 bp in length
150154 154046 contig of 3893 bp in length
154047 154146 contig of 100 bp in length
154147 156225 contig of 2079 bp in length
156226 156325 contig of 100 bp in length
156326 158578 contig of 2253 bp in length
158579 158678 contig of 100 bp in length
158679 161124 contig of 2446 bp in length
161125 161224 contig of 100 bp in length
161225 161224 contig of 100 bp in length

Sequence updated (26-May-2000).
* NOTE: This is a 'working draft' sequence. It currently
* consists of 32 contigs. The true order of the pieces
* is not known and their order in this sequence record is
* arbitrary. Gaps between the contigs are represented as
* runs of N, but the exact sizes of the gaps are unknown.
* This record will be updated with the finished sequence
* as soon as it is available and the accession number will
* be preserved.
1 23792: contig of 23792 bp in length
2 23892: gap of 100 bp
3 23893 45465: contig of 21573 bp in length
4 45466 45566: gap of 100 bp
5 45566 58167: contig of 12602 bp in length
6 58168 58267: gap of 100 bp
7 58268 71173: contig of 12906 bp in length
8 71174 71273: gap of 100 bp
9 71274 81253: contig of 9980 bp in length
10 81254 81354: gap of 100 bp
11 81354 91080: contig of 9727 bp in length
12 91081 91180: gap of 100 bp
13 91181 97387: contig of 6207 bp in length
14 97388 97487: gap of 100 bp
15 97488 104902: contig of 7415 bp in length
16 104903 105002: gap of 100 bp
17 105003 111107: contig of 6105 bp in length
18 111108 111207: gap of 100 bp
19 111208 116851: contig of 5644 bp in length
20 116852 116951: gap of 100 bp
21 116952 123585: contig of 6634 bp in length
22 123586 123685: gap of 100 bp
23 123686 128985: contig of 5300 bp in length
24 128986 129085: gap of 100 bp
25 129086 134233: contig of 5148 bp in length
26 134234 134334: gap of 100 bp
27 134334 138588: contig of 4255 bp in length
28 138589 138688: gap of 100 bp
29 138689 142573: contig of 3885 bp in length
30 142574 142673: gap of 100 bp
31 142674 146286: contig of 3613 bp in length
32 146287 146386: gap of 100 bp
33 146387 150053: contig of 3667 bp in length
34 150054 150153: gap of 100 bp
35 150154 154046: contig of 3893 bp in length
36 154047 154146: gap of 100 bp
37 154147 156225: contig of 2079 bp in length
38 156226 156325: gap of 100 bp
39 156326 158578: contig of 2253 bp in length
40 158579 158678: gap of 100 bp
41 158679 161124: contig of 2446 bp in length
42 161125 161224: gap of 100 bp

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* 161225 contig of 2335 bp in length
* 163559: gap of 100 bp
* 165360 contig of 1879 bp in length
* 165338: gap of 100 bp
* 165639 contig of 1311 bp in length
* 166949: gap of 100 bp
* 167049: gap of 100 bp
* 166950 contig of 1191 bp in length
* 168240: gap of 100 bp
* 168241: gap of 100 bp
* 168341 contig of 1239 bp in length
* 169579: gap of 100 bp
* 169678: gap of 100 bp
* 169680 contig of 1299 bp in length
* 170978: gap of 100 bp
* 171079 contig of 1002 bp in length
* 172080: gap of 100 bp
* 172081 contig of 1334 bp in length
* 173514: gap of 100 bp
* 173615 contig of 1000 bp in length
* 174615: gap of 100 bp
* 174715 contig of 1262 bp in length
* 175976: gap of 100 bp
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## FEATURES

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Location/Qualifiers

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/mol\_type="genomic DNA"

/db\_xref="taxon:9606"

/chromosome="18"

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154147..156225

Query Match

Best Local Similarity 95.5%; Score 478.6; DB 2; Length 177097;

Matches 487; Conservative 0; Mismatches 14; Indels 0; Gaps 0;

QY 1 CCACATGATTATCTCAATAGTCGAGAAAGCCATTTCACAAAATTTACAACTTTTCAT 60

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Db 147438 CCACATGATTATCTCAATAGTCGAGAAAGCCATTTCACAAAATTTCAACAACTTCAT 147497
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Db 147558 CTATGACAAACCACAGCCCAATATCATCTGAATGGGCAAAAATTTGGAAGCATTTCCCTTT 147617
QY 181 GAAAAAGGGGCAAGACAGAGGATGCCCTCTCTCACCACCTCTTCAACATAGTGTGGGA 240
Db 147618 GAAAACTGGGCAAGACAGAGGATGCCCTCTCTCACCACCTCTTCAACATAGTGTGGGA 147677
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Db 147738 GGAAGTCAAAATGTCCTGTTTGCAGATGATCATGATTGTATATCTAGAAAACCCATCGT 147797
QY 361 CTCAGCCCAAAATCTCCTTAAGCTGATAAGCACTTCAGCAAACTCTCAGATACAAAT 420
Db 147798 CTCAGCCCAAAATCTCCTTAAGCTGATAAGCACTTCAGCAAACTCTCAGATACAAAT 147857
QY 421 CAATGTACAAAATTCACAAAGCACTTTATATCATCAATAACAGACAAAACAGAGAGCCAAAT 480
Db 147858 CAATGTACAAAATTCACAAAGCACTTTATATCATCAATAACAGACAAAACAGAGAGCCAAAT 147917
QY 481 CATGAGTGAATCCCATTCAC 501
Db 147918 CATGAGTGAATCCCATTCAC 147938

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## RESULT 11

AC007052

LOCUS

DEFINITION

AC007052

AC007052.4

KEYWORDS

SOURCE

ORGANISM

REFERENCE

AUTHORS

TITLE

JOURNAL

REFERENCE

AUTHORS

179726 bp DNA linear PRI 23-MAR-1999  
Homo sapiens chromosome 18, clone hRPK.411\_H\_24, complete sequence.

AC007052

AC007052.4

HTG.

Homo sapiens (human)

Homo sapiens

Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;

Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.

1 (bases 1 to 179726)

Birren, B., Linton, L., Nusbaum, C. and Lander, E.

Homo sapiens chromosome 18, clone hRPK.411\_H\_24

Unpublished

2 (bases 1 to 179726)

Birren, B., Linton, L., Nusbaum, C., Lander, E., Allen, N., Anderson, M.,

Baker, J., Baldwin, J., Barna, N., Beckerly, R., Benn, J., Brown, A.,

Castle, A., Cerny, J., Colangelo, M., Collins, S., Collymore, A.,

Cooke, P., Dearellano, K., Depayre, E., Devon, K., Dewar, K.,

Donelan, L., Doyle, M., Ferreira, P., FitzHugh, W., Forrest, C.,

Funke, R., Gage, D., Galagan, J., Gardyna, S., Gilbert, D., Grant, G.,

Hagos, B., Heaford, A., Horton, L., Howland, J. C., Jones, C., Kann, L.,

Karatas, A., Lehoczy, J., Lieu, C., Locke, K., Macdonald, P.,

Marquis, N., McEwan, P., McGurk, A., McKernan, K., McLaughlin, J.,

Meldrum, J., Molla, M., Morris, W., Morrow, J., Mychaleckyj, J.,

Naylor, J., Niloff, M., O'Connor, T., O'Donnell, P., Pavlin, B.,

Peterson, K., Pollara, V., Riley, R., Roberts, D., Roy, A., Severy, P.,

Stange-Thomann, N., Stojanovic, N., Stone, C., Subramanian, A.,

Teefaye, S., Torruella-Miller, I., Vassiliev, H., Vo, A., Wagner, A.,

Wheeler, J., Wu, X., Wyman, D., Ye, W. J. and Zody, M.

Direct Submission

Submitted (09-MAR-1999) Whitehead Institute/MIT Center for Genome

Research, 320 Charles Street, Cambridge, MA 02141, USA

3 (bases 1 to 179726)

Birren, B., Linton, L., Nusbaum, C., Lander, E., Allen, N., Anderson, M.,

Baker, J., Baldwin, J., Barna, N., Beckerly, R., Benn, J., Brown, A.,

Castle, A., Cerny, J., Colangelo, M., Collins, S., Collymore, A.,

Cooke, P., Dearellano, K., Depayre, E., Devon, K., Dewar, K.,

Donelan, L., Doyle, M., Ferreira, P., FitzHugh, W., Forrest, C.,

Funke, R., Gage, D., Galagan, J., Gardyna, S., Gilbert, D., Grant, G.,

Hagos, B., Heaford, A., Horton, L., Howland, J. C., Jones, C., Kann, L.,

Karatas, A., Lehoczy, J., Lieu, C., Locke, K., Macdonald, P.,

Marquis, N., McEwan, P., McGurk, A., McKernan, K., McLaughlin, J.,

Meldrum, J., Molla, M., Morris, W., Morrow, J., Mychaleckyj, J.,

Naylor, J., Niloff, M., O'Connor, T., O'Donnell, P., Pavlin, B.,

Peterson, K., Pollara, V., Riley, R., Roberts, D., Roy, A., Severy, P.,

Stange-Thomann, N., Stojanovic, N., Stone, C., Subramanian, A.,

Teefaye, S., Torruella-Miller, I., Vassiliev, H., Vo, A., Wagner, A.,

Wheeler, J., Wu, X., Wyman, D., Ye, W. J. and Zody, M.

Direct Submission

Submitted (09-MAR-1999) Whitehead Institute/MIT Center for Genome

Research, 320 Charles Street, Cambridge, MA 02141, USA

3 (bases 1 to 179726)

Birren, B., Linton, L., Nusbaum, C., Lander, E., Allen, N., Anderson, M.,

Baker, J., Baldwin, J., Barna, N., Beckerly, R., Benn, J., Brown, A.,

Castle, A., Cerny, J., Colangelo, M., Collins, S., Collymore, A.,

Cooke, P., Dearellano, K., Depayre, E., Devon, K., Dewar, K.,

Donelan, L., Doyle, M., Ferreira, P., FitzHugh, W., Forrest, C.,

Funke, R., Gage, D., Galagan, J., Gardyna, S., Gilbert, D., Grant, G.,

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# TITLE

Submitted (23-MAR-1999) Whitehead Institute/MIT Center for Genome Research, 320 Charles Street, Cambridge, MA 02141, USA

On Mar 24, 1999 this sequence version replaced gi:4432872.

All repeats were identified using RepeatMasker: Smit, A.F.A. &

Green, P. (1996-1997)

<http://p1.genome.washington.edu/RM/RepeatMasker.html>.

## FEATURES

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* 32178 48716: contig of 16539 bp in length
* 48717 48816: gap of 100 bp
* 48817 68998: contig of 20182 bp in length
* 68999 103010: contig of 100 bp
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Db 136191 GAAACGGGCAACAGACAGGATGCCCTCTCTCACCCTCTCTTATTCACATAGTGTGGA 136250
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SOURCE Homo sapiens (human)
ORGANISM Homo sapiens
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE
1 (bases 1 to 186780)
AUTHORS Kimmerly, W., Bondoc, M., Cheng, J., Connolly, K. S., Gunning, K. M.,
Kadner, K., Miguel, T., Miller, C., Pitluck, S., Pollard, M.,
Rojeski, H., Subramanian, S. and Martin, C. H.
Rojeski, H., Subramanian, S. and Martin, C. H.
TITLE Sequencing of human chromosome 5
JOURNAL Unpublished
REFERENCE 2 (bases 1 to 186780)
AUTHORS Ricke, D. O.
TITLE Large Scale Sequence Analysis and Annotation with the Sequence
Comparison Analysis (SCAN) System
JOURNAL Unpublished
REFERENCE 3 (bases 1 to 186780)
AUTHORS Kimmerly, W., Bondoc, M., Cheng, J., Connolly, K. S., Gunning, K. M.,
Kadner, K., Miguel, T., Miller, C., Pitluck, S., Pollard, M.,
Rojeski, H., Subramanian, S. and Martin, C. H.
Direct Submission
TITLE Submitted (01-OCT-1998) Human Genome Center, DOE Joint Genome
JOURNAL Institute, Lawrence Berkeley National Laboratory, MS 74-157,
Berkeley, CA 94720, U.S.A.
COMMENT Sequence submitted by:
DOE Joint Genome Institute.
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SOURCE Homo sapiens
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Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
REFERENCE 1 (bases 1 to 196869)
AUTHORS Birren,B., Linton,L., Nusbaum,C., Lander,E., Allen,N., Anderson,S.,
Barna,N., Bastien,V., Boguslavsky,L., Bouckigalter,B., Brown,A.,
Canarata,J., Campopiano,A., Choepel,Y., Colangelo,M., Collins,S.,
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McPheeters,R., Meldrim,J., Meneus,L., Mihova,T., Mlenga,V.,

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Murphy, T., Naylor, J., Nguyen, C., Norbu, C., Norman, C.H., O'Connor, T., O'Donnell, P., O'Neill, D., Oliver, J., Peterson, K., Phunkhang, P., Pierre, N., Pollara, V., Raymond, C., Ratta, R., Rieback, M., Riley, R., Rise, C., Rogov, P., Roman, J., Rosetti, M., Roy, A., Santos, R., Schauer, S., Schuback, R., Seaman, S., Severy, P., Sougne, C., Spencer, B., Stange-Thomann, N., Stojanovic, N., Strause, N., Subramanian, A., Talamas, J., Testa, S., Theodore, J., Travers, M., Travis, N., Trigilio, J., Vassiliev, H., Viel, R., Vo, A., Wilson, B., Wu, X., Wyman, D., Ye, W., Young, G., Zainoun, J., Zembek, L., Zimmer, A. and Zody, M.

## TITLE

Direct Submission  
Submitted (06-JAN-2001) Whitehead Institute/MIT Center for Genome Research, 320 Charles Street, Cambridge, MA 02141, USA  
On Feb 25, 2001 this sequence version replaced gi:12043614.  
All repeats were identified using RepeatMasker:  
Smit, A.F.A. & Green, P. (1996-1997)  
http://ftp.genome.washington.edu/RM/RepeatMasker.html

## COMMENT

Center: Whitehead Institute/ MIT Center for Genome Research

Center code: WIBR

Web site: http://www-seq.wi.mit.edu

Contact: sequence\_submissions@genome.wi.mit.edu

----- Project Information

Center project name: L12339

Center clone name: 869\_L\_2

----- Summary Statistics

Sequencing vector: Plasmid; n/a; 100% of reads

Chemistry: Dye-terminator Big Dye; 100% of reads

Assembly program: Phrap; version 0.960731

Consensus quality: 189512 bases at least Q40

Consensus quality: 193548 bases at least Q30

Consensus quality: 194822 bases at least Q20

Insert size: 198000; agarose-fp

Insert size: 195669; sum-of-contigs

Quality coverage: 5.4 in Q20 bases; agarose-fp

Quality coverage: 5.5 in Q20 bases; sum-of-contigs

----- NOTE: This is a 'working draft' sequence. It currently consists of 13 contigs. The true order of the pieces is not known and their order in this sequence record is arbitrary. Gaps between the contigs are represented as runs of N, but the exact sizes of the gaps are unknown. This record will be updated with the finished sequence as soon as it is available and the accession number will be preserved.

\* 11489: contig of 11489 bp in length

\* 11589: gap of 100 bp

\* 11590: 12389: contig of 800 bp in length

\* 12390: 12489: gap of 100 bp

\* 12490: 15301: contig of 2812 bp in length

\* 15302: 15401: gap of 100 bp

\* 15402: 17297: contig of 1896 bp in length

\* 17298: 17397: gap of 100 bp

\* 17398: 22220: contig of 4823 bp in length

\* 22221: 22220: gap of 100 bp

\* 22321: 27991: contig of 5671 bp in length

\* 27992: 28091: gap of 100 bp

\* 28092: 31242: contig of 3151 bp in length

\* 31243: 31342: gap of 100 bp

\* 31343: 41247: contig of 9905 bp in length

\* 41248: 41347: gap of 100 bp

\* 41348: 49982: contig of 8635 bp in length

\* 49983: 50082: gap of 100 bp

\* 50083: 64395: contig of 14313 bp in length

\* 64396: 64495: gap of 100 bp

\* 64496: 79369: contig of 14874 bp in length

\* 79370: 79469: gap of 100 bp

\* 79470: 151119: contig of 71650 bp in length

\* 151120: 151219: gap of 100 bp

\* 151220: 196869: contig of 45650 bp in length.

Location/Qualifiers

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Qy 121 CTATGACAACTCAGCCCAATATCATACTGAATGGGCAAAACTGGAAGCATTCCTTT 180  
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ACCESSION AC011401  
VERSION AC011401.9 GI:23396212  
KEYWORDS HTG.  
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ORGANISM Homo sapiens  
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;  
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.  
REFERENCE 1 (bases 1 to 199275)  
AUTHORS DOE Joint Genome Institute and Stanford Human Genome Center.  
TITLE Direct Submission  
JOURNAL Unpublished  
REFERENCE 2 (bases 1 to 199275)  
AUTHORS DOE Joint Genome Institute.  
TITLE Direct Submission  
JOURNAL Submitted (06-OCT-1999) Production Sequencing Facility, DOE Joint Genome Institute, 2800 Mitchell Drive, Walnut Creek, CA 94598, USA  
REFERENCE 3 (bases 1 to 199275)  
AUTHORS DOE Joint Genome Institute.  
TITLE Direct Submission  
JOURNAL Submitted (07-MAR-2002) Production Sequencing Facility, DOE Joint Genome Institute, 2800 Mitchell Drive, Walnut Creek, CA 94598, USA  
REFERENCE 4 (bases 1 to 199275)  
AUTHORS DOE Joint Genome Institute and Stanford Human Genome Center.  
TITLE Direct Submission  
JOURNAL Submitted (01-OCT-2002) DOE Joint Genome Institute, 2800 Mitchell Drive, Walnut Creek, CA 94598, USA  
COMMENT On Oct 1, 2002 this sequence version replaced gi:12830130.  
Draft Sequence Produced by DOE Joint Genome Institute  
www.jgi.doe.gov  
Finishing Completed at Stanford Human Genome Center  
www.sngc.stanford.edu  
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Best Local Similarity 97.2%; Pred. No. 2e-108;  
Matches 487; Conservative 0; Mismatches 14; Indels 0; Gaps 0;

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Job time : 1969.4 secs



GenCore version 5.1.6  
Copyright (c) 1993 - 2003 Compugen Ltd.

OM nucleic - nucleic search, using sw model

Run on: November 6, 2003, 21:13:51 ; Search time 310.835 Seconds  
(without alignments)  
8684.478 Million cell updates/sec

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Scoring table: IDENTITY\_NUC  
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Searched: 2552756 seqs, 1349719017 residues

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Post-processing: Minimum Match 0%  
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Listing first 45 summaries

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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4	311.2	31.1	26997	22	AA546747
5	296	29.6	15418	21	AAA63785
6	296	29.6	15418	24	AA54997
7	296	29.6	15418	24	AA138595
8	296	29.6	15418	24	AA138601

9	293.6	29.4	27067	25	AA50021	Human secreted pro
10	288	28.8	30393	22	AAK67239	Human immune/haema
c 11	286.6	28.7	44147	24	ABK84481	Human cDNA differe
12	280.8	28.1	32206	22	AAK89374	Human digestive sy
13	270.4	27.0	20420	22	AAK73165	Human immune/haema
14	270.4	27.0	20420	24	ABK69933	Human secreted pro
15	267.4	26.7	19965	22	AAK73166	Human immune/haema
16	267.4	26.7	19965	24	ABK69932	Human secreted pro
c 17	266.4	26.6	6971	24	ABL33237	Human immune syste
18	263	26.3	165199	24	ABK83460	Human cDNA differe
c 19	263	26.3	227968	24	ABK83497	Human cDNA differe
c 20	261.4	26.1	12069	24	ABK39930	Human chemically p
c 21	260.6	26.1	20420	22	AAK73165	Human immune/haema
c 22	260.6	26.1	20420	24	ABK69933	Human secreted pro
23	258.4	25.8	82938	24	ABV72623	Human transporter
c 24	258	25.8	19965	22	AAK73166	Human immune/haema
c 25	258	25.8	19965	24	ABK69932	Human secreted pro
c 26	257.6	25.8	12069	24	ABK39930	Human chemically p
c 27	256	25.6	870	14	AAQ39248	Sequence of cosmid
c 28	253.6	25.4	80595	20	AAV83939	HC-contig derived
c 29	253	25.3	106746	21	AAA10225	Human PCNA-1 genom
c 30	252.6	25.3	80240	20	AAV83940	NC-contig derived
c 31	252.6	25.3	82938	24	ABV72623	Human transporter
c 32	250	25.0	21537	24	ABL33999	Human immune syste
c 33	247.4	24.7	121600	24	ABT10748	Human breast cance
c 34	245	24.5	6286	22	AA54591	Tumour suppressor
c 35	242	24.2	1691080	24	ABX08336	Human phosphodiester
c 36	240.4	24.0	13573	24	ABK33869	Human immune syste
c 37	240.2	24.0	165199	24	ABK83460	Human cDNA differe
c 38	237.4	23.7	40023	24	ABL51954	Human solute carri
c 39	235.2	23.5	21537	24	ABL33998	Human immune syste
c 40	234.2	23.4	26997	22	AA546748	Tumour suppressor
c 41	234.2	23.4	80595	20	AAV83939	HC-contig derived
c 42	234.2	23.4	154902	24	ABQ8198	Human osteoblast d
c 43	232.8	23.3	110000	22	AA54800	Nucleotide sequenc
c 44	232.2	23.2	80240	20	AAV83940	NC-contig derived
c 45	226.8	22.7	15577	19	AAV35616	SHOX gene prelimin

ALIGNMENTS

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AC	
DT	18-JUN-2002 (first entry)
XX	Genomic nucleotide sequence encoding human RGS-4 protein.
XX	RGS-4; schizophrenia; human; regulator of G protein signalling 4;
KW	neuroleptic; polymorphism; schizotypy; schizoid; chromosome 1q21-22;
KW	gene; ds.
OS	Homo sapiens.
XX	
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FT	variation
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XX 24-AUG-2001; 2001WO-US26622.
XX
XX 24-AUG-2000; 2000US-228021P.
XX
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```
PA (UYPI-) UNIV PITTSBURGH.
XX
XX Levitt PR, Mirnics K, Kodavali VC, Nimgaonkar VL;
XX
XX WPI; 2002-292070/33.
XX
XX Diagnosing, assessing susceptibility and treating schizophrenia,
XX involves observing regulator of G-protein signalling 4, RGS4 levels in a
XX subject -
XX
XX Claim 1; Page 20-33; 112pp; English.
XX
XX This invention relates to a novel method for diagnosing schizophrenia
XX or determining susceptibility to schizophrenia in a human. The method
XX comprises obtaining from a patient a DNA sample and detecting variations
XX in the regulator of G-protein signalling 4 (RGS4) gene. Alternatively,
XX the method involves measuring RGS4 mRNA or protein levels in a tissue
XX sample from the patient and determining if there is a reduced level.
XX The method of the invention is useful for diagnosing and determining
XX susceptibility to schizophrenia. The invention also comprises a method
XX that is useful for treating schizophrenia which includes a prophylactic
XX treatment. The method of genotyping polymorphic variants in the RGS-4
XX gene is applied to diagnosing pathologies of the schizophrenic spectrum,
XX such as in particular schizotypy, schizoid individuals, etc. This
XX method offers the possibility of diagnosing schizophrenia by a
XX biological test and no longer exclusively by clinical evaluations.
XX The present sequence represents the genomic DNA encoding the human
XX regulator of G-protein signalling 4 (RGS4) protein used in the method of
XX the invention. The gene for the RGS4 protein is located on human
XX chromosome 1q21-22.
XX
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XX Best Local Similarity 100.0%; Pred. No. 5.5e-150;
XX Matches 1000; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
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Db 481 TATATATATATACACATATATATATATATATATATATATATATATATATATATAC 540
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AAS46748/c
ID AAS46748 standard; DNA; 26997 BP.
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AC AAS46748;
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DT 18-DEC-2001 (first entry)
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DE Tumour suppressor gene derived chemically modified sequence #472.
XX
KW Human; tumour suppressor gene; oncogene; antitumour; cytostatic;
KW cancer; tumour; CpG dinucleotide; single-nucleotide polymorphism; SNP;
KW cytosine methylation; ds.
XX
OS Homo sapiens.
XX
PN WO200168912-A2.
XX
PD 20-SEP-2001.
XX
PF 15-MAR-2001; 2001WO-EP02955.
XX
PR 15-MAR-2000; 2000DE-1013847.
PR 06-APR-2000; 2000DE-1019058.
PR 07-APR-2000; 2000DE-1019173.
PR 30-JUN-2000; 2000DE-1032529.
PR 01-SEP-2000; 2000DE-1043826.
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FA (EPIG-) EPIGENOMICS AG.
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PI Olek A, Piepenbrock C, Berlin K;
XX
DR WPI; 2001-602752/68.
XX
PT Fragments of chemically modified genes associated with tumour suppressor
PT genes and oncogenes, useful in designing primers and probes for
PT analysing diseases associated with cytosine methylation state e.g.
PT cancer -
XX
PS Claim 1; SEQ ID No 472; 27pp; English.
XX
CC The invention relates to a nucleic acid comprising a sequence of 18
CC bases, of a segment of chemically pretreated DNA (CP DNA) e.g. with
CC bisulphite, of genes associated with tumour suppression and
CC oncogenes having a sequence taken from 536 (actually 533 since
CC numbers 408, 458 and 500 are missing from the sequence listing) sequences
CC (Ss) and sequences complementary to (Ss). The nucleic acid may be a
CC peptide nucleic acid-oligomer (PNA) of at least 9 nucleotides and may
CC form part of a set of probes for detecting the cytosine methylation state
CC and/or single nucleotide polymorphisms and also to be used in an
CC array for analysing diseases associated with CpG dinucleotides e.g.
CC cancers and tumours. The probes can also be used in a method for
CC ascertaining genetic and/or epigenetic parameters for the diagnosis
CC and/or therapy of existing diseases or the predisposition to specific
CC diseases, by analysing cytosine methylations. The parameters may be
CC compared to another set of genetic and/or epigenetic parameters, the
CC differences serving as basis for diagnosis and/or prognosis events which
CC are disadvantageous to patients. The present sequence is one of the
CC 533 genomic sequences derived from tumour suppressor genes and
CC oncogenes. Sequences with even numbered Seq ID numbers are the
CC complementary sequence of the corresponding odd numbered sequence (e.g.
CC ID 2 and ID1, ID 536 and ID 535, except for those whose partner sequence
CC is missing).
CC Note: The sequence data for this patent did not form part
CC of the printed specification, but was obtained in electronic
CC format directly from WIPO at
CC ftp.wipo.int/pub/published_pct_sequences.
XX
SQ Sequence 26997 BP; 7650 A; 354 C; 5924 G; 13069 T; 0 other;
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Best Local Similarity 66.0%; Pred. No. 8.2e-43;
Matches 498; Conservative 0; Mismatches 253; Indels 4; Gaps 2;

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[illegible]

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XX	
XX	
AC	ABSS54997;
XX	
XX	
DT	10-DEC-2002 (first entry)
XX	
XX	
DE	Lambda clone containing human TERT genomic insert.
XX	
KW	Telomerase reverse transcriptase; TERT; replication-conditional virus;
KW	adenovirus replication gene; cancer cell; lung; pancreatic cancer;
KW	medulloblastoma; cervical carcinoma; fibrosarcoma; osteosarcoma;
KW	cytology; replication defective adenovirus vector; congenital defect;
KW	proinflammatory; antiinflammatory; heterologous effector gene;
KW	cancer therapy; cytostatic; gene therapy; lambda clone; human; ds.
XX	
OS	Bacteriophage lambda.
OS	Homo sapiens.
OS	Synthetic.
XX	
XX	

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PN	WO200253760-A2.	
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XX	11-JUL-2002.	
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XX	17-DEC-2001; 2001WO-US48785.	
PF		
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XX	18-DEC-2000; 2000US-256418P.	
PR		
XX	(GERO-) GERON CORP.	
PA		
XX		
PI	Irving JM, Lebkowski JS;	
XX		
DR	WPI; 2002-723123/78.	
XX		
XX	Novel replication-conditional virus useful for cytolysis of target	
PT	cells e.g. cancer cells and preparing a medicament for treating cancer,	
PT	comprises heterologous replication element in an adenovirus-based	
FT	construct	
PT		
XX		
XX	Claim 11; Page 26-29; 32pp; English.	
PS		
XX		
CC	The present invention relates to a new replication-conditional virus with	
CC	a genome comprising adenovirus replication genes and one or more	
CC	heterologous gene(s) that functionally replaces one or more adenovirus	
CC	gene(s) required for replication or assembly of the virus. The invention	
CC	is useful for killing a cancer cell (such as lung, pancreatic cancer,	
CC	medulloblastoma, cervical carcinoma, fibrosarcoma or osteosarcoma),	
CC	killing a cell expressing TERT (telomerase reverse transcriptase), and in	
CC	preparing a medicament for treating cancer and a condition associated	
CC	with increased expression of TERT in affected cells, in a subject.	
CC	The invention is also useful for cytolysis of specific target cells.	
CC	The invention is further useful for producing replication defective	
CC	adenovirus vector which is useful for transient expression of a	
CC	heterologous therapeutic gene to correct a congenital defect, introducing	
CC	proinflammatory or antiinflammatory activity, enhancing telomerase	
CC	function, and delivering heterologous effector genes that induce killing	
CC	of the transduced cells. The invention is more safe for use in cancer	
CC	therapy. The present nucleic acid sequence represents the human TERT	
CC	sequence contained within a lambda clone sequence of the invention.	
XX		
XX	Sequence 15418 BP; 4518 A; 3797 C; 3765 G; 3338 T; 0 other;	

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Qy	60	AGACAGGCATGGTGATACAGCCCTGTATCCACGACTCTTCGGAGGCCGAGCAGAGAA	119		
Db	706	AGCTGGGTGTGTGACACATGCCTGTATCCAGGTACTCAGAGGCTTAAGCAGAGAA	765		
Qy	120	TCACTTGAAACCTGCTGGGGGTGGAGGTTGGGGGAGCAAGATCATGCCATTGCATCTCCAG	179		
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DT 05-NOV-2001 (first entry)  
XX Human digestive system antigen genomic sequence SEQ ID NO: 2950.  
DE Human; digestive system antigen; gene therapy; cancer; appendicitis;  
XX ulcerative colitis; infection; Hirschsprung's disease; chronic colitis;  
KW digestive system disorder; Meckel's diverticulum; ds.  
KW  
XX  
XX Homo sapiens.  
XX WO200155314-A2.  
XX  
XX 02-AUG-2001.  
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XX 17-JAN-2001; 2001WO-US01324.  
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XX 31-JAN-2000; 2000US-0179065.  
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KW Human; immune; haematopoietic; immune/haematopoietic antigen; cancer;  
KM cytotatic; gene therapy; vaccine; metastasis; ds.  
XX  
OS Homo sapiens.  
XX WO200157182-A2.  
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OM nucleic - nucleic search, using sw model

Run on: November 6, 2003, 21:13:51; Search time 621.98 Seconds  
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Title: US-09-939-209a-3\_COPY\_3000\_5000  
Perfect score: 2001  
Sequence: 1 attgtataattagcactc.....cagttcttctaaagcctatt 2001

Scoring table: IDENTITY\_NUC  
Gapop 10.0, Gapext 1.0

Searched: 2552756 seqs, 1349719017 residues  
Total number of hits satisfying chosen parameters: 5105512

Minimum DB seq length: 0  
Maximum DB seq length: 2000000000  
Post-processing: Minimum Match 0%  
Maximum Match 100%  
Listing first 45 summaries

Database :			
N Geneseq_19Jun03.*			
1:	/SIDSI/gcgdata/geneseq/geneseq-emb1/NA1980.DAT.*		
2:	/SIDSI/gcgdata/geneseq/geneseq-emb1/NA1981.DAT.*		
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24:	/SIDSI/gcgdata/geneseq/geneseq-emb1/NA2002.DAT.*		
25:	/SIDSI/gcgdata/geneseq/geneseq-emb1/NA2003.DAT.*		
SUMMARIES			
Result No.	Score	Query Match Length DB ID	Description
1	2001	100.0 20300 24	ABK47337 Genomic nucleotide
2	173.8	8.7 126512 24	ABN83429 Human transporter
3	170	8.5 965 22	AAH33769 Human colon cancer
4	167.4	8.4 168575 22	AAH21613 Human hypocretin r
5	164.2	8.2 541 25	ABZ71922 Human cancer-assoc
6	164.2	8.2 462586 25	ABQ84281 Chromosome 2 Asthm
7	161.6	8.1 1503841 24	ABT00010 Human neuregulin 1
8	161.6	8.1 1503841 24	ABT01503 Human neuregulin 1

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

ALIGNMENTS			
RESULT 1			
ABK47337	ABK47337 standard; DNA; 20300 BP.		
ID	ABK47337	standard; DNA; 20300 BP.	
XX	AC	ABK47337;	
XX	DT	18-JUN-2002 (first entry)	
XX	DE	Genomic nucleotide sequence encoding human RGS-4 protein.	
XX	KW	RGS-4; schizophrenia; human; regulator of G protein signalling 4;	
KW	KW	neuroleptic; polymorphism; schizotypy; schizoid; chromosome 1q21-22;	
XX	XX	Gene; ds.	
OS	OS	Homo sapiens.	
XX	XX	Key	Location/Qualifiers
FT	FT	variation	replace (4121,T)
FT	FT	variation	/*tag= a
FT	FT	variation	/standard name= "Single-nucleotide polymorphism"
FT	FT	variation	replace (4123,A)
FT	FT	variation	/*tag= b
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FT	FT	variation	replace (4790,T)
FT	FT	variation	/*tag= e

Human neuregulin-1  
Human neuregulin-1  
Human osteoblast d  
Human immune/haema  
Human secreted pro  
Human inflammatory  
cDNA encoding a pr  
Human immune/haema  
Human PACAP genom  
Human cDNA encodin  
Human cDNA differe  
Human immune/haema  
Human prostate exp  
Human rTS-alpha ge  
Human rTS-beta gen  
MAGE-B cluster DNA  
Human genomic DNA  
Human genomic DNA  
Gene #2225 used to  
Stomach cancer rel  
Nucleotide sequenc  
Human pancreatic c  
Human digestive sy  
Human chloride int  
Human nervous syst  
Human methionine a  
Human phosphodiester  
Human pancreatic c  
Human digestive sy  
Phosphoribosyl gly  
Human secreted pro  
Human ABC transpor  
Human endocrine po  
Human immune/haema  
Human osteoblast d  
Human voltage-acti



Qy	541	ACTGTTGTGAAGTGCACCAATGACTATATATTTTCCCTATATACTTGTGATATTTTGTGCACTCG	600
Db	3540	ACTGTTGTGAAGTGCACCAATGACTATATATTTTCCCTATATACTTGTGATATTTTGTGCACTCG	3599
Qy	601	CCCATGAGAAATGTAGTCTAAGATCAAAAGGATGCAAGAAATGGGTCTCTATCCAGTATAGTAC	660
Db	3600	CCCATGAGAAATGTAGTCTAAGATCAAAAGGATGCAAGAAATGGGTCTCTATCCAGTATAGTAC	3659
Qy	661	CCACTACACTGGTGGATGTCAATATGTATTTGTTAGATTAATATCTCTCAAGAAATGACGACC	720
Db	3660	CCACTACACTGGTGGATGTCAATATGTATTTGTTAGATTAATATCTCTCAAGAAATGACGACC	3719
Qy	721	TTTCTCAGACACATAAAAAGATGCTCAATATAATAAAAGTTTGTGTGAACCTGAACTATTGGCA	780
Db	3720	TTTCTCAGACACATAAAAAGATGCTCAATATAATAAAAGTTTGTGTGAACCTGAACTATTGGCA	3779
Qy	781	AATGTACATCATCGGATTTTAAAGAGAGCGAAACAGAGTCTGGCTCAAAACACCATACT	840
Db	3780	AATGTACATCATCGGATTTTAAAGAGAGCGAAACAGAGTCTGGCTCAAAACACCATACT	3839
Qy	841	TCTAGAGTGCAATAAGAGGTAGCAGTCTGAATTACCACTGGCGACAGAGCAAAAAAGAGCTTG	900
Db	3840	TCTAGAGTGCAATAAGAGGTAGCAGTCTGAATTACCACTGGCGACAGAGCAAAAAAGAGCTTG	3899
Qy	901	ACCACGGGTACTGTGAAGACATTTTCAGGTTGTATGGCCACAGAAACAGGGGAAATACATAAA	960
Db	3900	ACCACGGGTACTGTGAAGACATTTTCAGGTTGTATGGCCACAGAAACAGGGGAAATACATAAA	3959
Qy	961	TGTGTGGGAATATTCAGTGGTCTGGGATGACTACATAGTAGAATATATTAATGAAGAAAGAG	1020
Db	3960	TGTGTGGGAATATTCAGTGGTCTGGGATGACTACATAGTAGAATATATTAATGAAGAAAGAG	4019
Qy	1021	TGGAAGGGAAGATGAAAAGTTGGAATGGGGATGAAATTATGAAAGTACCAGAAATGTTATG	1080
Db	4020	TGGAAGGGAAGATGAAAAGTTGGAATGGGGATGAAATTATGAAAGTACCAGAAATGTTATG	4079
Qy	1081	CTAAGGAATCTAGATTTTAAATGTGAAGGGCAAAATTGAAGTCTCTGGGCACGTTTACAAAAC	1140
Db	4080	CTAAGGAATCTAGATTTTAAATGTGAAGGGCAAAATTGAAGTCTCTGGGCACGTTTACAAAAC	4139
Qy	1141	TAGAGGTCTATAAGTTTACCCCTAATTTACCAAGNTTTCCTAGAGATCTATATTTGGAAAT	1200
Db	4140	TAGAGGTCTATAAGTTTACCCCTAATTTACCAAGNTTTCCTAGAGATCTATATTTGGAAAT	4199
Qy	1201	CCAGATCTGCTCTCTGTAAAGTTTCAAGCACTTTTCCATGACACACATCTGTTCTTTTCCA	1260
Db	4200	CCAGATCTGCTCTCTGTAAAGTTTCAAGCACTTTTCCATGACACACATCTGTTCTTTTCCA	4259
Qy	1261	CTTGCACAATGCAAAATGAACTCTTATGAAACTGCTGTTTCTATCTCTGGGCTAAATGTTGC	1320
Db	4260	CTTGCACAATGCAAAATGAACTCTTATGAAACTGCTGTTTCTATCTCTGGGCTAAATGTTGC	4319
Qy	1321	AGAAAAAGATTTAATCTTTGGGATAGGCTATTTTGGGTTTTTCTCTACTTCTTGGGAA	1380
Db	4320	AGAAAAAGATTTAATCTTTGGGATAGGCTATTTTGGGTTTTTCTCTACTTCTTGGGAA	4379
Qy	1381	ACAAGGTTTTCTTCCCCTGGCTAATTAAGTGTGGTATTGTTCTTCCAGGGAAATCAGTGA	1440
Db	4380	ACAAGGTTTTCTTCCCCTGGCTAATTAAGTGTGGTATTGTTCTTCCAGGGAAATCAGTGA	4439
Qy	1441	TGCATCACCTGCTGCTATCAAAATGTCAGGGTGGAGTTCCTGATTTTATGTCATGTGCCA	1500
Db	4440	TGCATCACCTGCTGCTATCAAAATGTCAGGGTGGAGTTCCTGATTTTATGTCATGTGCCA	4499
Qy	1501	CAAGCTTGGTGCAAGAAATTTGGACATTTCCCAAAGTAAGACATATCTGGGAAATCCC	1560
Db	4500	CAAGCTTGGTGCAAGAAATTTGGACATTTCCCAAAGTAAGACATATCTGGGAAATCCC	4559
Qy	1561	TGTTTACTCTTCTGGTATACAGATCTCTCAGGCCCATATCTTTTGTCTTTTATGCTCTAAA	1620
Db	4560	TGTTTACTCTTCTGGTATACAGATCTCTCAGGCCCATATCTTTTGTCTTTTATGCTCTAAA	4619
Qy	1621	AATCAATPAACTGAACCTCTCATGTGATGCTTAGGCCATTTGTAGTAAACAATAAAGAGGAGG	1680

Db	4620	AATCAATAA	CTGAACTCTCA	TTGATGCTAGGCCA	TTGTAGTAAACAATTAAGAGGAGG	4679
Qy	1681	GAGGCTTCTG	CAACTGAGAGGA	AAATGTCATCTG	GAAGTGGTGCACGACAGCCTGGGGC	1740
Db	4680	GAGGCTTCTG	CAACTGAGAGGA	AAATGTCATCTG	GAAGTGGTGCACGACAGCCTGGGGC	4739
Qy	1741	TGAGCCTTGG	CTACATCTCTG	CCCAAGTCGAGG	ATCAGTGCCCAATTTAAACATCTGGTAG	1800
Db	4740	TGAGCCTTGG	CTACATCTCTG	CCCAAGTCGAGG	ATCAGTGCCCAATTTAAACATCTGGTAG	4799
Qy	1801	AACATAAGAA	CGCAACGCCCTG	CCCAATGACTTA	TTTCCTGCAATTTGATACCGTCAATC	1860
Db	4800	AACATAAGAA	CGCAACGCCCTG	CCCAATGACTTA	TTTCCTGCAATTTGATACCGTCAATC	4859
Qy	1861	CTTGAGAAAT	GTTTTCTTTT	TGTTCTCCCTG	AGCAAGGTTGGAAAAATTTGAAATTTACC	1920
Db	4860	CTTGAGAAAT	GTTTTCTTTT	TGTTCTCCCTG	AGCAAGGTTGGAAAAATTTGAAATTTACC	4919
Qy	1921	TAGAGACACA	CATAGTTTCAT	CTCTCTGTGGCT	GGAATGTCGTGCCCCCAGTAGGAA	1980
Db	4920	TAGAGACACA	CATAGTTTCAT	CTCTCTGTGGCT	GGAATGTCGTGCCCCCAGTAGGAA	4979
Qy	1981	ACAGTTCTT	CTAAAGCCTATT	2001		
Db	4980	ACAGTTCTT	CTAAAGCCTATT	5000		
RESULT 2						
ABN83429						
ID	ABN83429 standard; DNA; 126512 BP.					
XX						
AC	ABN83429;					
DT	21-AUG-2002 (first entry)					
DE	Human transporter protein gene.					
XX						
KW	Human; sodium/calcium exchanger; transporter; brain; heart; kidney; lung;					
KW	spleen; testis; leukocyte; foetal brain; chromosome 14; gene;					
KW	single nucleotide polymorphism; SNP; ds.					
XX						
OS	Homo sapiens.					
EH	Key	Location/Qualifiers				
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FT		replace(741..742,C-)				
FT		/*tag= b				
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FT		/note= "This variation is an indel"				
FT	variation	replace(2002,T)				
FT		/*tag= c				
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FT	CDS	2010..124505				
FT		/*tag= d				
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FT		/note= "Contains 5 introns"				
FT	exon	2010..3793				
FT		/*tag= e				
FT		/number= 1				
FT	variation	replace(2381,C)				
FT		/*tag= f				
FT		/standard name= "Single nucleotide polymorphism"				
FT	intron	3794..109509				
FT		/*tag= g				
FT		/number= 1				
FT	variation	replace(5165,T)				
FT		/*tag= h				
FT		/standard name= "Single nucleotide polymorphism"				
FT	variation	replace(5402,G)				
FT		/*tag= i				



FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(6794,C)  
FT /tag= j  
FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(9883,G)  
FT /tag= k  
FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(10210,C)  
FT /tag= l  
FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(12220,G)  
FT /tag= m  
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FT replace(13842,G)  
FT /tag= n  
FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(14200,A)  
FT /tag= o  
FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(15878,T)  
FT /tag= p  
FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(16030,G)  
FT /tag= q  
FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(16292,C)  
FT /tag= r  
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FT replace(16506,G)  
FT /tag= s  
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FT /tag= z  
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FT /standard\_name= "Single nucleotide polymorphism"  
FT /note= "This variation is an indel"  
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FT /tag= az  
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FT /tag= ba  
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FT replace(69280,T)  
FT /tag= bd  
FT /standard\_name= "Single nucleotide polymorphism"  
FT replace(70647,T)  
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FT

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FT variation replace(71867,T)
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FT /standard_name= "Single nucleotide polymorphism"
FT variation replace(71900,T)
FT /*tag= bg
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FT variation replace(71901,A)
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FT variation replace(72369,T)
FT /*tag= bi

Query Match      8.7%; Score 173.8; DB 24; Length 136512;
Best Local Similarity 83.4%; Pred. No. 3.5e-34;
Matches 221; Conservative 0; Mismatches 42; Indels 2; Gaps 2;

QY 308 TACCTTTCTAGTTAGATAG-TTATGATACACAAATATATTTTCATTGTGTATAAATTT 366
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
57055 TGCCTTTTCTAGGCTAGATATGTTTATGATACACATACATACATTTTACCATTTGTTCCTAATG 57114

QY 367 CCTACAGTATTCAGTACATGCTGCTGATACAGTTTGTACCTAGGAGTAATAGGCTA 426
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
57115 CCTACAGTTTCCAGTACATGCTGCTGATACAGTTTGTACCTAGGAGCAATAGGCTA 57174

QY 427 TACCATACAGCTTAGGTTGTAGTAGGCTATAACCATCTAGGTTTGTCTAAGTACATTTCT 486
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
57175 TACCATACAGCTTAGGTTGTAGTAGGCTAT-ACCCTTAGGCTGGGTAAGTACATCTCT 57233

QY 487 ATGATATTTCCCAATGATGAAATCCATCACTACATTTTCTCAGAAATGTTTTCATGTT 546
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
57234 ATGATGTTTTCACAGTGTAGAACTTCTCTAATGACAAATTTCTCAGAAATGATCCAGTT 57293

QY 547 GTGAAGTGACCCATGACTATATTTT 571
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
57294 GTTAAGTGAGCATGACAGTACTAT 57318

RESULT 3
AAH33769/c
ID AAH33769 standard; cDNA; 965 BP.
XX
XX AAH33769;
AC
XX
DT 03-SEP-2001 (first entry)
DE Human colon cancer antigen encoding cDNA SEQ ID NO:825.
XX
XX Human; colon cancer; colon cancer antigen; diagnosis; detection;
XX colorectal carcinoma; ss.
XX
XX Homo sapiens.
XX
XX WO200122920-A2.
XX
XX 05-APR-2001.
XX
XX 28-SEP-2000; 2000WO-US26524.
XX
XX 29-SEP-1999; 99US-0157137.
XX
XX 03-NOV-1999; 99US-0163280.
XX
XX (HUMA-) HUMAN GENOME SCI INC.
XX
XX Ruben SM, Barash SC, Birse CE, Rosen CA;
XX
XX WPI; 2001-235357/24.
XX
XX P-PSDB; AAG74338.
XX
XX Nucleic acids encoding 4277 human colon cancer-associated polypeptides,
XX useful for preventing, diagnosing and/or treating colorectal cancers -
XX
XX Claim 1; Page 2780-2781; 9803pp; English.
XX

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CC AAH32943 to AAH37195 and AAG73514 to AAG77788 represent human colon
CC cancer-associated nucleic acid molecules (N) and proteins (P), where
CC the proteins are collectively known as colon cancer antigens. The colon
CC cancer antigens have cytostatic activity and can be used in gene
CC therapy and vaccine cytoaction. N and P may be used in the prevention,
CC diagnosis and treatment of diseases associated with inappropriate P
CC expression. For example, N and P may be used to treat disorders
CC associated with decreased expression by rectifying mutations or deletions
CC in a patient's genome that affect the activity of P by expressing
CC inactive proteins or to supplement the patient's own production of P.
CC Additionally, N may be used to produce the colon cancer-associated Pe
CC by inserting the nucleic acids into a host cell and culturing the cell
CC to express the proteins. N and P can be used in the prevention, diagnosis
CC and treatment of colorectal carcinomas and cancers. AAH37196 to AAH37204
CC and AAG77789 represent sequences used in the exemplification of the
CC present invention.
CC N.B. Pages 666 to 682 and page 7053 of the sequence listing were
CC missing at time of publication, meaning no sequences are present for
CC SEQ ID NO:1027 to 1052, 7921 and 7922.
XX
SQ Sequence 965 BP; 315 A; 126 C; 174 G; 349 T; 1 other;

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Query Match      8.5%; Score 170; DB 22; Length 965;
Best Local Similarity 72.2%; Pred. No. 5e-34;
Matches 283; Conservative 0; Mismatches 90; Indels 19; Gaps 4;

QY 182 ATGTGCCCAVACAATGTTTTCAGTCAGGAGTACAGCAAAATGTATCTGGCCCAATAT 241
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
631 ATACGACATACAGTCATGTTTCGATCAACAATTGACCACATATGACAGAGATCCTATAG 572

QY 242 ATTATA-----AGCTGAGAAATTTCTATTAACCTAGTGATATGCGAGCCATCAAG--TGT 295
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
571 ATTATAATGGAACCTGAAATAATTCCTATCACCTAGTGATGCCACAGCCATTGTTACATTGT 512

QY 296 AATCGAGGACATTACCTTTTCTATGTTTATAGTATGTTAGATACACAAATATTTTCATTG 355
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
511 AGCAATATGTTATTTCTGTTCTATGTTTATGATTACAGATAC-----CATTTG 464

QY 356 TGTATATATTTCTACAGTATTTTCAGTACAGTAACATGCTGTACAGGTTTGTAACTAGAA 415
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
463 TGTACAGTTTGCCTACAGTATTTCAATACAGTACCATGCTGTACAGGCTGTAGCCAGAA 404

QY 416 GTAATAGGCTATACCATACAGCTTAGGTTGTGTATAGGCTATAACCATCTAGGTTTGTGT 475
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
403 GCGACAGACTTTGCCATCTAGCCAGGTGTGCTAGGCTCT-ACCACCTAGGTTTGTGT 345

QY 476 AAGTACATTCTATGATATTTCCCAATGATGAAATCACCTAACTACACATTTCTCAGAAAT 535
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
344 AAGTACATTCTATGATTTTACACATGATGAAATCACCTAACTACACATTTCTCAGAAAT 285

QY 536 GTTTCACCTGTTGTGAAGTGACCCATGACTATA 567
DB ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
284 GTAGCTCTGTCATGAAGTACACATTTACTGTA 253

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RESULT 4
AAH21613
ID AAH21613 standard; DNA; 168575 BP.
XX
XX AAH21613;
AC
XX
DT 10-AUG-2001 (first entry)
XX
XX Human hypocretin receptor 2 (HCRTR2) gene SEQ ID NO:1.
XX
XX Human; narcolepsy; hypocretin receptor 2; orexin receptor 2; HCRTR2;
XX diagnosis; ds.
XX
XX Homo sapiens.
XX
XX WO200130991-A2.
XX
XX 03-MAY-2001.
XX

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XX 22-AUG-2000; 2000WO-US23021.
XX PF
XX 25-OCT-1999; 99US-0426290.
XX PR
XX (DECO-) DECODE GENETICS EHF.
XX PA
XX Olafsdottir BR, Gulcher J;
XX PI
XX WPI; 2001-300504/31.
XX DR
XX P-PSDB; AAB98007.
XX DR
XX
XX Gene for hypocretin (orexin) receptor 2 (HCRTR2) which is associated
XX with narcolepsy, useful in methods of diagnosis of narcolepsy and
XX PT pharmaceutical compositions for therapy -
XX PT
XX Claim 1; Fig 1; 85pp; English.
XX PS
XX
XX The present invention describes the human hypocretin (orexin) receptor 2
XX (HCRTR2) gene (given in AAH21613), which is associated with narcolepsy.
XX CC Identification of the HCRTR2 nucleic acid molecule permits the diagnosis
XX CC of narcolepsy. A method from the present invention is provided for
XX CC treating narcolepsy by administering to the individual an isolated
XX CC HCRTR2 nucleic acid in a therapeutically effective amount so that the
XX CC cells produce native HCRTR2 receptor. The diagnosis of narcolepsy has
XX CC been difficult to differentiate from other conditions such as chronic
XX CC fatigue syndrome or other sleep disorders but detection of HCRTR2
XX CC nucleic acid makes it possible to accurately diagnose narcolepsy.
XX CC AAH21541 to AAH21612 represent primers used in the identification of the
XX CC narcolepsy gene in an example from the present invention. AAH21613
XX CC represents the HCRTR2 gene which encodes the HCRTR2 protein given in
XX CC AAB98007.
XX CC
XX SQ Sequence 168575 BP; 55308 A; 29672 C; 29838 G; 53757 T; 0 other;
Query Match 8.4%; Score 167.4; DB 22; Length 168575;
Best Local Similarity 79.2%; Pred. No. 1.8e-32;
Matches 224; Conservative 0; Mismatches 56; Indels 3; Gaps 2;
QY 311 CTTTCTATGTTAGATATGTT-AGATACACAAATATATTCATTGTGTTATATTTTCT 369
DB 57581 CTTTCTATGTTAGATATGTTAGATACACAAATGCTTATCATTTGTGTTATATTTGCT 57640
QY 370 ACAGTATTCAGTACAGTACATGCTGACAGGTTTGTAACTAGGAGTATAGGCTATAC 429
DB 57641 ACAGTGTTCAGTACAGTACATGCTGACAGGTTTATAGGCTAGGAGCAATTGGCTATAC 57700
QY 430 CATACAGCTTAGGTGTAGTGTAGGCTATACCATCTAGGTTTGTGTAAGTACATTCATG 489
DB 57701 CCTATAGCCTAGGTGTGTAGTGTAGGCTATA--CCATTAGATTTGTGTAAGCATACCCCTATG 57758
QY 490 ATATTCACCAATGATGAATACCTTAACATACACATTTCTCAGATGTTTTCACGTGTGTG 549
DB 57759 ATGTTTGCACAATGATGAATACCTTAAGGATGCAATTTCTCAGCATATATCCAGTCATT 57818
QY 550 AAGTGACCCCATGACTATATTTTCTATATATCTTATATCTTGTATTTTGT 592
DB 57819 AAGCAAGACTGACTATATTTATTAGGCTATTTTATTCTATAG 57861
RESULT 5
ABZ71922/c
ID ABZ71922 standard; cDNA; 541 BP.
XX
AC ABZ71922;
XX
XX 01-APR-2003 (first entry)
XX
XX Human cancer-associated gene SEQ ID NO 229.
XX
XX Human; cancer; stomach cancer; cytostatic; gene; ss.
XX
XX Homo sapiens.
OS
XX
XX WO200283899-A1.
XX PN
XX 24-OCT-2002.
XX PD
XX
XX 28-MAR-2002; 2002WO-JP03038.
XX PF
XX
XX 10-APR-2001; 2001JP-0112039.
XX PR
XX 21-SEP-2001; 2001JP-0290193.
XX PR
XX (TAKA-) TAKARA BIO INC.
XX PA
XX
XX Yoshikawa Y, Okamoto S, Oura T, Mineno J, Asada K, Kato I;
XX PI Inoue H, Mori M;
XX PI
XX WPI; 2003-093022/08.
XX DR
XX
XX Measuring changes in expression of 264 cancer associated genes for
XX PT detection of stomach cancer and screening of potential anticancer
XX PT agents -
XX PT
XX Claim 1; Page 235-236; 266pp; Japanese.
XX PS
XX
XX The invention relates to a method for the detection of cancer in which a
XX CC change in the expression of 1 or more of 264 specified cancer associated
XX CC genes, ABZ71694-ABZ71957, or of sequences at least 80% homologous to them
XX CC in the specimen tissue as compared to normal tissue is observed. The
XX CC genes are used in detection, diagnosis and treatment of cancer.
XX CC especially of stomach cancer. The present sequence is that of a cancer
XX CC associated polynucleotide of the invention.
XX CC
XX SQ Sequence 541 BP; 166 A; 110 C; 117 G; 148 T; 0 other;
Query Match 8.2%; Score 164.2; DB 25; Length 541;
Best Local Similarity 81.1%; Pred. No. 1.3e-32;
Matches 215; Conservative 0; Mismatches 48; Indels 2; Gaps 2;
QY 304 ACATTACCTTTTCTATGTTTAGATAT-GTTAGATACACAAATATATTTTCATTGTGTTATA 362
DB 395 ACCTTACCTTTTCTATGTTTAGCTCTGGGTAGATAGACAAATACTTACCATTGTGTTTA 336
QY 363 ATTTCCTACAGTATTCAGTACAGTACATGCTGTACAGGTTTGTAACTAGGAGTAAATAG 422
DB 335 ATTGCGCTGCAATATTCAGTACAGTACATGCTGTACAGGCTCGTAGGCTAGGACCAACAG 276
QY 423 GCTATACCAATACAGCTTAGGTGTAGTGTAGGCTATACCATCTAGGTTGTGTAAGTACA 482
DB 275 GCTATACCAATACAGCTTAGGTGTAGTGTAGGCTAT-ACCATCTAGGTTGTGTAAGTACA 217
QY 483 TTCTATGATATTCCTCAATGATGAATCACTAACTACACATTTCTCAGAATGTTTTCAC 542
DB 216 CTCTTTGATATTTGCACACCTTGAATTCGCCGAATGACACATTTCTCAGGACATATTC 157
QY 543 TGTGTGTGAGTGTGACCCATGACTATA 567
DB 156 TGTCCACCAAGCAACCCCATGACTGTA 132
RESULT 6
ABQ84281/c
ID ABQ84281 standard; DNA; 462586 BP.
XX
XX ABQ84281;
XX AC
XX
XX 20-FEB-2003 (first entry)
XX DT
XX
XX Chromosome 2 Asthma Locus DNA sequence.
XX DE
XX
XX DPP10; dipeptidyl peptidase; prololigopeptidase; enzyme; asthma;
XX KW antiinflammatory; antiasthmatic; antipsoriatic; antiarthritic;
XX KW antirheumatic; vaccine; gene therapy; inflammatory disease;
XX KW inflammatory bowel disease; atopy; rheumatoid arthritis; psoriasis;
XX KW chromosome 2q14; gene; ds.
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Query Match 8.1%; Score 161.6; DB 24; Length 1503841;  
Best local similarity 82.5%; Pred. No. 1.4e-30;  
Matches 221; Conservative 0; Mismatches 44; Indels 3; Gaps 3;

QY 304 ACATTACCTTTCTATGTTTAGATATG-TTAGATACACAATATATTTCATTGTTGTTATA 362  
|||||







diagnosis and treatment of diseases associated with inappropriate NR1G1 expression. For example, they may be used to treat disorders associated with decreased expression by rectifying mutations or deletions in a patient's genome that affect the activity of NR1G1 by expressing of inactive proteins or to supplement the patients own production of NR1G1. Additionally, the gene may be used to produce NR1G1 polypeptides, by inserting the nucleic acids into a host cell and culturing the cell to express the protein. The gene may also be used as DNA probes and primers in diagnostic assays to detect and quantitate the presence of similar nucleic acids in samples, and therefore which patients may be in need of restorative therapy. The NR1G1 polypeptides may also be used as antigens in the production of antibodies against NR1G1 and in assays to identify modulators of NR1G1 expression and activity. Anti-NR1G1 antibodies and antagonists may also be used to down regulate expression and activity. Anti-NR1G1 antibodies may also be used as diagnostic agents for detecting the presence of NR1G1 polypeptides in samples. NR1G1 is associated with schizophrenia which may be prevented, diagnosed and/or treated by the above methods.

Sequence 1503900 BP; 452487 A; 281874 C; 288074 G; 480092 T; 1373 other;

Query Match 8.1%; Score 161.6; DB 22; Length 1503900;  
Best Local Similarity 82.5%; Pred. No. 1.4e-30;  
Matches 221; Conservative 0; Mismatches 44; Indels 3; Gaps 3;

QY 304 ACATTACCTTTCTATGTTAGATATG-TTAGATACACAAATATATTCATTGCTTATA 362  
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QY 363 ATTTCTACAGTATTCAGTACAGTAACATGCTGTACAGGTTTGTAACTAGGAGTAATAG 422  
DB |||||||  
QY 73826 ACTGTACCTTTCTATGTTAGTATGTTAGTATTCACAAATATTCATTGTTGTA 73885  
DB |||||||  
QY 363 ATTTCTACAGTATTCAGTACAGTAACATGCTGTACAGGTTTGTAACTAGGAGTAATAG 422  
DB |||||||  
QY 73886 ATTGCTACAGTATTCAGTACAGTAACATGCTGTACAGGTTTGTAGCCTAGGACATAG 73945  
DB |||||||  
QY 423 GCTATACCATACAGCTTAGGTGTGTAGTACGCTATACCATCTAGGTTTGTAGTACA 482  
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QY 73946 ACTCTACCATATAGCCTACATCTGCAATAGGCTA-CACTATCTAGGTTTGTAGTACA 74004  
DB |||||||  
QY 483 TTCTATGATATTTCCACAAATGATGAATCACTACTACATATTCCTCAGATGTTTCCAC 542  
DB |||||||  
QY 74005 CTCTATGATGTTTCCACCAATGAGNAATCACCCGA-GACATTTCTCAGAACGTATCCC 74063  
DB |||||||  
QY 543 TGTGTGAAGTGACCCACTGACTATATTT 570  
DB |||||||  
QY 74064 TGTCAATTAAGTGAACATGACTGTAGTT 74091  
DB |||||||

RESULT 10  
AAK96733  
XX AAK96733 standard; DNA; 1503900 BP.  
XX AC AAK96733;  
XX DT 17-DEC-2001 (first entry)  
XX DE Human neuregulin-1 gene.  
XX KW Human; neuregulin 1 gene; schizophrenia; gene therapy; ds.  
XX OS Homo sapiens.  
XX PN W0200164877-A2.  
XX PD 07-SEP-2001.  
XX PF 28-FEB-2001; 2001WO-US06377.  
XX PR 28-FEB-2000; 2000US-0515716.  
XX PA (DECO-) DECODE GENETICS EHF.  
XX PI Stefansson H, Steinhorsdottir V, Gulcher JR;  
XX WPI; 2001-514841/56.

DR P-PSDB; AAG67938, AAG67939, AAG67940, AAG67941, AAG67942, AAG67943,  
DR AAG67944, AAG67945, AAG67946, AAG67947, AAG67948, AAG67949,  
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DR AAG67962, AAG67963, AAG67964, AAG67965, AAG67966, AAG67967,  
DR AAG67968, AAG67969, AAG67970, AAG67971, AAG67972, AAG67973,  
DR AAG67974, AAG67975.  
XX  
PT Neuregulin 1 nucleic acids and proteins useful for diagnosing  
PT preventing and treating schizophrenia -  
XX  
PS Disclosure; Page 345-756; 756pp; English.  
XX  
XX This sequence represents the human neuregulin 1 gene of the invention.  
CC The invention also relates to fragments or variants of the neuregulin 1  
CC gene. The gene and its proteins may be used in the prevention, diagnosis  
CC and treatment of diseases associated with inappropriate neuregulin 1  
CC expression, such as schizophrenia. For example they may be used to treat  
CC disorders associated with decreased neuregulin 1 expression by rectifying  
CC mutations or deletions in a patient's genome that affect the activity of  
CC neuregulin 1 by expressing inactive proteins or to supplement the  
CC patients own production of polypeptides. Additionally, the gene may be  
CC used to produce the neuregulin 1 protein, by inserting the nucleic acids  
CC into a host cell and culturing the cell to express the protein. The gene  
CC and its complementary sequences may also be used as DNA probes in  
CC diagnostic assays to detect and quantitate the presence of similar  
CC nucleic acids in samples, and therefore which patients may be in need of  
CC restorative therapy. The protein may also be used as antigens in the  
CC production of antibodies against neuregulin 1 and in assays to identify  
CC modulators of neuregulin 1 expression and activity. The antibodies and  
CC antagonists may also be used to down regulate expression and activity.  
CC The antibodies may also be used as diagnostic agents for detecting the  
CC presence of neuregulin 1 in samples.  
XX  
SQ Sequence 1503900 BP; 452487 A; 281874 C; 288074 G; 480092 T; 1373 other;  
Query Match 8.1%; Score 161.6; DB 22; Length 1503900;  
Best Local Similarity 82.5%; Pred. No. 1.4e-30;  
Matches 221; Conservative 0; Mismatches 44; Indels 3; Gaps 3;

QY 304 ACATTACCTTTCTATGTTAGATATG-TTAGATACACAAATATATTCATTGCTTATA 362  
DB |||||||  
QY 73826 ACTGTACCTTTCTATGTTAGTATGTTAGTATTCACAAATATTCATTGTTGTA 73885  
DB |||||||  
QY 363 ATTTCTACAGTATTCAGTACAGTAACATGCTGTACAGGTTTGTAACTAGGAGTAATAG 422  
DB |||||||  
QY 73886 ATTGCTACAGTATTCAGTACAGTAACATGCTGTACAGGTTTGTAGCCTAGGACATAG 73945  
DB |||||||  
QY 423 GCTATACCATACAGCTTAGGTGTGTAGTACGCTATACCATCTAGGTTTGTAGTACA 482  
DB |||||||  
QY 73946 ACTCTACCATATAGCCTACATCTGCAATAGGCTA-CACTATCTAGGTTTGTAGTACA 74004  
DB |||||||  
QY 483 TTCTATGATATTTCCACAAATGATGAATCACTACTACATATTCCTCAGATGTTTCCAC 542  
DB |||||||  
QY 74005 CTCTATGATGTTTCCACCAATGAGNAATCACCCGA-GACATTTCTCAGAACGTATCCC 74063  
DB |||||||  
QY 543 TGTGTGAAGTGACCCACTGACTATATTT 570  
DB |||||||  
QY 74064 TGTCAATTAAGTGAACATGACTGTAGTT 74091  
DB |||||||

RESULT 11  
ABQ88198/c  
ID ABQ88198 standard; cDNA; 154902 BP.  
XX  
XX ABQ88198;  
XX AC  
XX DT 18-SEP-2002 (first entry)  
XX DE Human osteoblast differentiation related cDNA SEQ ID NO 105.  
XX KW Human; osteoblast; stem cell differentiation; bone tissue deposition;  
XX KW osteoporosis; osteopathic; ss.

```
XX OS Homo sapiens.
XX PN WO200250301-A2.
XX XX
XX PD 27-JUN-2002.
XX PF
XX PF 18-DEC-2001; 2001WO-US48276.
XX PR 18-DEC-2000; 2000US-255882P.
XX PR 24-APR-2001; 2001US-285691P.
XX XX
XX PA (GENE-) GENE LOGIC INC.
XX PA (PROC ) PROCTER & GAMBLE CO.
XX XX
XX PI Ji D, Axelrod DW, Cook JS, Jaiswal N, Einstein R, Houghton A;
XX PI Mertz L;
XX XX
XX DR WPI; 2002-557663/59.
XX XX
XX PT Use of genes and their expression profiles associated with osteoblast
XX PT differentiation for screening modulators bone formation, for diagnosing
XX PT or treating e.g. osteoporosis, or as markers for the differentiation
XX PT process -
XX XX
XX PS Claim 1; SEQ ID NO 105; 78bp + Sequence Listing; English.
XX XX
XX CC The invention relates to genes and their expression profiles are used
XX CC for:
XX CC (a) screening modulators of precursor stem cell differentiation into
XX CC osteoblasts, or bone tissue deposition;
XX CC (b) diagnosing abnormal deposition of bone tissue, abnormal rate of
XX CC osteoblast formation or osteoporosis; or
XX CC (c) treating or monitoring treatment of the conditions cited in (b), or
XX CC monitoring the progression of bone tissue deposition.
XX CC Specific conditions include postmenopausal osteoporosis, glucocorticoid
XX CC osteoporosis or male osteoporosis, osteopenia, osteodystrophy,
XX CC drug-induced abnormalities in bone formation or bone loss, conditions
XX CC that involve altered bone metabolism (e.g. idiopathic juvenile
XX CC osteoporosis), skeletal disease linked to breast cancer, mastocytosis,
XX CC Fanconi syndrome or fibrous dysplasia. The present sequence is that of an
XX CC osteoblast differentiation associated cDNA marker of the invention.
XX CC Note: The sequence data for this patent did not form part of the printed
XX CC specification, but was obtained in electronic format directly from WIPO
XX CC at ftp.wipo.int/pub/published_pct_sequences.
XX SQ
XX SQ Sequence 154902 BP; 43917 A; 31458 C; 32848 G; 46679 T; 0 other;
XX
XX Query Match 8.1%; Score 161.2; DB 24; Length 154902;
XX Best Local Similarity 80.9%; Pred. No. 7.2e-31;
XX Matches 212; Conservative 0; Mismatches 48; Indels 2; Gaps 2;
XX
XX QY 308 TACCTTTTCATGTTAGATATG-TTAGATACACAAATATATTTTCATTGTTATAATTT 366
XX Db 138160 TACCTTTTCAATATTTAGATATGTTAGAAATACAGATACCTTACCAITGTTACAATTG 138101
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XX QY 367 CCTACAGATATTCAGTACGTAACATGCTGTACAGGTTTGTAACTAGAGTAATAGCCTA 426
XX Db 138100 CCTACAGCATGCGATCAGTAACATGCTGTACAGGTTTGTAGCCTGGAGTAATTTGCTA 138041
XX
XX QY 427 TACCATACAGCTTAGGTGTGTAGTAGGCTATAACCATCTAGGTTTGTAGTACATTTCT 486
XX Db 138040 CCCCATATAGCTTAGGTGTGTAGAGGTTAT-ACCATCTAGGTTTGTAGGCTACTCT 137982
XX
XX QY 487 ATGATATTCACCAATGATGAAATCACTCACTACACATTTCTCAGAAATGTTTCACATGTT 546
XX Db 137981 ATAAATGTTTATACAAATGACAAAATCTCTCAACACACATTTTTCAGAACGTCCTCCTTT 137922
XX
XX QY 547 GTGAGATGACCCACGACTATAT 568
XX Db 137921 GTTAAGCGCACATGACTACAT 137900
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RESULT 12
AAK62341/c
ID AAK62341 standard; cDNA; 402 BP.
XX XX
XX AC AAK62341;
XX XX
XX DT 06-NOV-2001 (first entry)
XX XX
XX DE Human immune/haematopoietic antigen encoding cDNA SEQ ID NO:7401.
XX XX
XX KW Human; immune; haematopoietic; immune/haematopoietic antigen; cancer;
XX KW cytostatic; gene therapy; vaccine; metastasis; ss.
XX XX
XX OS Homo sapiens.
XX XX
XX PN WO200157182-A2.
XX XX
XX PD 09-AUG-2001.
XX XX
XX PF 17-JAN-2001; 2001WO-US01354.
XX XX
XX PF 31-JAN-2000; 2000US-0179065.
XX PR 04-FEB-2000; 2000US-0180628.
XX PR 24-FEB-2000; 2000US-0184664.
XX PR 02-MAR-2000; 2000US-0186350.
XX PR 16-MAR-2000; 2000US-0189874.
XX PR 17-MAR-2000; 2000US-0190076.
XX PR 18-APR-2000; 2000US-0198123.
XX PR 19-MAY-2000; 2000US-0205515.
XX PR 07-JUN-2000; 2000US-0209467.
XX PR 28-JUN-2000; 2000US-0214886.
XX PR 30-JUN-2000; 2000US-0215135.
XX PR 07-JUL-2000; 2000US-0216647.
XX PR 07-JUL-2000; 2000US-0216880.
XX PR 11-JUL-2000; 2000US-0217487.
XX PR 11-JUL-2000; 2000US-0217496.
XX PR 14-JUL-2000; 2000US-0218290.
XX PR 26-JUL-2000; 2000US-0220963.
XX PR 26-JUL-2000; 2000US-0220964.
XX PR 14-AUG-2000; 2000US-0224518.
XX PR 14-AUG-2000; 2000US-0224519.
XX PR 14-AUG-2000; 2000US-0225213.
XX PR 14-AUG-2000; 2000US-0225214.
XX PR 14-AUG-2000; 2000US-0225266.
XX PR 14-AUG-2000; 2000US-0225267.
XX PR 14-AUG-2000; 2000US-0225268.
XX PR 14-AUG-2000; 2000US-0225270.
XX PR 14-AUG-2000; 2000US-0225447.
XX PR 14-AUG-2000; 2000US-0225757.
XX PR 14-AUG-2000; 2000US-0225758.
XX PR 14-AUG-2000; 2000US-0225759.
XX PR 18-AUG-2000; 2000US-0226279.
XX PR 22-AUG-2000; 2000US-0226681.
XX PR 22-AUG-2000; 2000US-0226688.
XX PR 22-AUG-2000; 2000US-0227182.
XX PR 23-AUG-2000; 2000US-0227009.
XX PR 30-AUG-2000; 2000US-0228924.
XX PR 01-SEP-2000; 2000US-0229287.
XX PR 01-SEP-2000; 2000US-0229343.
XX PR 01-SEP-2000; 2000US-0229344.
XX PR 01-SEP-2000; 2000US-0229345.
XX PR 03-SEP-2000; 2000US-0229509.
XX PR 05-SEP-2000; 2000US-0229513.
XX PR 06-SEP-2000; 2000US-0230437.
XX PR 06-SEP-2000; 2000US-0230438.
XX PR 08-SEP-2000; 2000US-0231242.
XX PR 08-SEP-2000; 2000US-0231243.
XX PR 08-SEP-2000; 2000US-0231244.
XX PR 08-SEP-2000; 2000US-0231413.
XX PR 08-SEP-2000; 2000US-0231414.
XX PR 08-SEP-2000; 2000US-0232080.
XX PR 08-SEP-2000; 2000US-0232081.
XX PR 12-SEP-2000; 2000US-0231968.
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21: /SIDSI/gcgdata/geneseq/geneseqn-emb1/NA2000.DAT.\*  
22: /SIDSI/gcgdata/geneseq/geneseqn-emb1/NA2001A.DAT.\*  
23: /SIDSI/gcgdata/geneseq/geneseqn-emb1/NA2001B.DAT.\*  
24: /SIDSI/gcgdata/geneseq/geneseqn-emb1/NA2002.DAT.\*  
25: /SIDSI/gcgdata/geneseq/geneseqn-emb1/NA2003.DAT.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

# SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	501	100.0	20300	24	ABK47337
2	180.6	36.0	1238	22	ABL26766
3	177.4	35.4	3165	25	ABX63113
4	45.4	9.1	10020	24	ABL34293
5	41.2	8.2	8845	22	AA546544
6	40.6	8.1	1456	24	ABQ61218
7	40.2	8.0	5661	24	ABQ67088
8	40	8.0	306	24	ABN94869

9	40	8.0	306	24	ABL63962	Breast cancer rela
10	39.8	7.9	9646	24	ABL33688	Human immune syste
11	39.6	7.9	2360	21	AAC66096	PSS3 cDNA sequence
12	39.6	7.9	2494	21	AAC66090	Rice sucrose synth
13	38.6	7.7	648	22	AAK69902	Human immune/haema
14	38.2	7.6	15732	22	AAV54123	Rabbit membrane pr
15	38.2	7.6	15732	22	AAS45388	Chemically pretrea
16	38.2	7.6	15732	22	ABK28233	DNA transcription
17	38	7.6	140167	24	ABT10146	Human breast cance
18	37.8	7.5	851	20	AAK95566	Nucleic acid seque
19	37.8	7.5	8372	23	ABL23336	Drosophila melanog
20	37.8	7.5	11473	24	ABK40030	Human immune syste
21	37.8	7.5	11473	24	ABL33355	C. albicans BAX-as
22	37.2	7.4	513	24	ABQ76618	Chemically treated
23	37.2	7.4	6228	24	ABL70469	Human gene regulat
24	37.2	7.4	6228	24	AAS61430	Signal transductio
25	37.2	7.4	112132	24	ABK31496	Human ATP-dependen
26	37.2	7.4	112132	24	ABK90888	Signal transductio
27	37	7.4	7508	24	ABL31206	Human immune syste
28	37	7.4	9964	24	ABL32099	Human immune syste
29	37	7.4	10048	24	ABQ67015	S. epidermidis ope
30	36.8	7.3	543	22	AAH53661	S. epidermidis gen
31	36.8	7.3	3666	22	AAH54866	Human brain expres
32	36.6	7.3	487	22	AAK18805	Human bone marrow
33	36.6	7.3	487	22	AAK44744	Probe #19407 used
34	36.6	7.3	487	22	AAI50721	Human liver single
35	36.6	7.3	487	23	ABK44404	Human genome-deriv
36	36.6	7.3	487	24	ABK18983	Human brain expres
37	36.6	7.3	491	22	AAK06024	Human bone marrow
38	36.6	7.3	491	22	AAK13760	Probe #6232 used t
39	36.6	7.3	491	22	AAK137546	Human liver single
40	36.6	7.3	491	23	ABK31354	Human genome-deriv
41	36.6	7.3	491	24	ABK06426	N. meningitidis pa
42	36.6	7.3	853	21	AAA81913	Arabidopsis profil
43	36.6	7.3	978	21	AAC64462	Arabidopsis profil
44	36.6	7.3	1162	21	AAC64461	Arabidopsis profil
45	36.6	7.3	1388	21	AAC64460	Arabidopsis profil

## ALIGNMENTS

### RESULT 1

ABK47337  
ID ABK47337 standard; DNA; 20300 BP.

XX AC ABK47337;

XX DT 18-JUN-2002 (first entry)

XX DE Genomic nucleotide sequence encoding human RGS-4 protein.

XX KW RGS-4; schizophrenia; human; regulator of G protein signalling 4;  
neuroleptic; polymorphism; schizotypy; schizoid; chromosome 1q21-22;  
KW gene; ds.

XX OS Homo sapiens.

XX FH Key

FT variation

FT replace (4121,T)

FT /tag= a

FT /standard name= "Single-nucleotide polymorphism"

FT replace (4123,A)

FT /tag= b

FT /standard name= "Single-nucleotide polymorphism"

FT replace (4368,C)

FT /tag= c

FT /standard name= "Single-nucleotide polymorphism"

FT replace (4621,C)

FT /tag= d

FT /standard name= "Single-nucleotide polymorphism"

FT replace (4790,T)

FT /tag= e

FT /standard\_name= "Single-nucleotide polymorphism"  
FT replace (4816,T)  
FT /\*tag= f  
FT /standard\_name= "Single-nucleotide polymorphism"  
FT replace (4970,T)  
FT /\*tag= g  
FT /standard\_name= "Single-nucleotide polymorphism"  
FT replace (5055,G)  
FT /\*tag= h  
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FT /\*tag= i  
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FT replace (5695,A)  
FT /\*tag= j  
FT /standard\_name= "Single-nucleotide polymorphism"  
FT replace (7375,T)  
FT /\*tag= k  
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FT replace (7759,A)  
FT /\*tag= l  
FT /standard\_name= "Single-nucleotide polymorphism"  
FT replace (8596,A)  
FT /\*tag= m  
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FT replace (9602..9610,CA)  
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FT replace (9892,A)  
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FT replace (11056,C)  
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FT /\*tag= v  
FT /standard\_name= "Single-nucleotide polymorphism"  
FT replace (12145,T)  
FT /\*tag= w  
FT /standard\_name= "Single-nucleotide polymorphism"  
FT replace (14367,G)  
FT /\*tag= x  
FT /standard\_name= "Single-nucleotide polymorphism"  
FT replace (17027..17029,GA)  
FT /\*tag= y  
FT /note= "Deletion polymorphism"  
FT replace (17630,T)  
FT /\*tag= z  
FT /standard\_name= "Single-nucleotide polymorphism"  
XX WO200216653-A2.  
XX  
XX 28-FEB-2002.  
XX  
XX 24-AUG-2001; 2001WO-US26622.  
XX  
XX 24-AUG-2000; 2000US-228021P.  
XX  
XX

PA (UYPI-) UNIV PITTSBURGH.  
XX  
XX Levitt PR, Mirnics K, Kodavali VC, Nimgaonkar VL;  
XX  
XX WPI; 2002-292070/33.  
XX  
XX Diagnosing, assessing susceptibility and treating schizophrenia,  
XX involves observing regulator of G-protein signalling 4, RGS4 levels in a  
XX subject -  
XX  
XX Claim 1; Page 20-33; 112pp; English.  
XX  
XX This invention relates to a novel method for diagnosing schizophrenia  
XX or determining susceptibility to schizophrenia in a human. The method  
XX comprises obtaining from a patient a DNA sample and detecting variations  
XX in the regulator of G-protein signalling 4 (RGS4) gene. Alternatively,  
XX the method involves measuring RGS4 mRNA or protein levels in a tissue  
XX sample from the patient and determining if there is a reduced level.  
XX The method of the invention is useful for diagnosing and determining  
XX susceptibility to schizophrenia. The invention also comprises a method  
XX that is useful for treating schizophrenia which includes a prophylactic  
XX treatment. The method of genotyping polymorphic variants in the RGS-4  
XX gene is applied to diagnosing pathologies of the schizophrenic spectrum,  
XX such as in particular schizotypy, schizoid individuals, etc. This  
XX method offers the possibility of diagnosing schizophrenia by a  
XX biological test and no longer exclusively by clinical evaluations.  
XX The present sequence represents the genomic DNA encoding the human  
XX regulator of G-protein signalling 4 (RGS4) protein used in the method of  
XX the invention. The gene for the RGS4 protein is located on human  
XX chromosome 1q21-22.  
XX  
XX SQ Sequence 20300 BP; 6157 A; 4102 C; 3775 G; 6266 T; 0 other;  
XX  
XX Query Match 100.0%; Score 501; DB 24; Length 20300;  
XX Best Local Similarity 100.0%; Pred. No. 1.7e-132;  
XX Matches 501; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
XX  
XX Qy 1 GACCATGTATAATGATGCTTCTTAATCCAAAGAGGAAAGGCATTGGAGTCAGTCCT 60  
XX Db 10000 GACCATGTATAATGATGCTTCTTAATCCAAAGAGGAAAGGCATTGGAGTCAGTCCT 10059  
XX  
XX Qy 61 AAGTAAGCTCCAGAAATTCCTGCTGCTGCTATTTCTTCCTCCAGGAAGCACTTCCTGATATT 120  
XX Db 10060 AAGTAAGCTCCAGAAATTCCTGCTGCTGCTATTTCTTCCTCCAGGAAGCACTTCCTGATATT 10119  
XX  
XX Qy 121 TTTTCTTACAGGCATATGATATAAACTATATTTTGCAGCATTTGACACTTTTTTTCCT 180  
XX Db 10120 TTTTCTTACAGGCATATGATATAAACTATATTTTGCAGCATTTGACACTTTTTTTCCT 10179  
XX  
XX Qy 181 TTTCTAGAAATTTAAACCTCTGACATTTGGTGGAGACATTGAGTACATTTTCCCATAT 240  
XX Db 10180 TTTCTAGAAATTTAAACCTCTGACATTTGGTGGAGACATTGAGTACATTTTCCCATAT 10239  
XX  
XX Qy 241 CCCTACATTTTCAGAGGATTTTCTGCTCGTTCACATTTACATTTGCTGATCGTCAGTCT 300  
XX Db 10240 CCCTACATTTTCAGAGGATTTTCTGCTCGTTCACATTTGCTGATCGTCAGTCT 10299  
XX  
XX Qy 301 TTTCTTCTCATCTCTTTTCAGGGGCTGGAGAGGAGGAGGAGACAGAGGAGCTGGTACTG 360  
XX Db 10300 TTTCTTCTCATCTCTTTTCAGGGGCTGGAGAGGAGGAGGAGACAGAGGAGCTGGTACTG 10359  
XX  
XX Qy 361 CAGAGCGGTGCTGTGATTTGGCTGGACGGTCTGAGCTGGCTATATAAGAGACCCCTACAG 420  
XX Db 10360 CAGAGCGGTGCTGTGATTTGGCTGGACGGTCTGAGCTGGCTATATAAGAGACCCCTACAG 10419  
XX  
XX Qy 421 GCTTAGCAGGAAGACGCTCAGAGGATTTCTGACATATCTTTACGGAGAGAGGCAAGT 480  
XX Db 10420 GCTTAGCAGGAAGACGCTCAGAGGATTTCTGACATATCTTTACGGAGAGAGGCAAGT 10479  
XX  
XX 481 ACGCTCAAAGCCGAAGCCACA 501  
XX  
XX 10480 ACGCTCAAAGCCGAAGCCACA 10500  
XX











PR 25-SEP-2000; 2000US-234923P.  
PR 25-SEP-2000; 2000US-234924P.  
PR 25-SEP-2000; 2000US-235077P.  
PR 25-SEP-2000; 2000US-235082P.  
PR 25-SEP-2000; 2000US-235134P.  
PR 25-SEP-2000; 2000US-235280P.  
PR 26-SEP-2000; 2000US-235637P.  
PR 26-SEP-2000; 2000US-235638P.  
PR 27-SEP-2000; 2000US-235711P.  
PR 27-SEP-2000; 2000US-235720P.  
PR 27-SEP-2000; 2000US-235840P.  
PR 27-SEP-2000; 2000US-235863P.  
PR 28-SEP-2000; 2000US-236028P.  
PR 28-SEP-2000; 2000US-236032P.  
PR 28-SEP-2000; 2000US-236033P.  
PR 28-SEP-2000; 2000US-236034P.  
PR 28-SEP-2000; 2000US-236109P.  
PR 28-SEP-2000; 2000US-236111P.  
PR 29-SEP-2000; 2000US-236942P.  
PR 29-SEP-2000; 2000US-236891P.  
PR 02-OCT-2000; 2000US-237172P.  
PR 02-OCT-2000; 2000US-237173P.  
PR 02-OCT-2000; 2000US-237278P.  
PR 02-OCT-2000; 2000US-237279P.  
PR 02-OCT-2000; 2000US-237295P.  
PR 02-OCT-2000; 2000US-237316P.  
PR 03-OCT-2000; 2000US-237425P.  
PR 03-OCT-2000; 2000US-237598P.  
PR 03-OCT-2000; 2000US-237604P.  
PR 03-OCT-2000; 2000US-237606P.  
PR 03-OCT-2000; 2000US-237608P.  
PR 01-NOV-2000; 2000US-244867P.  
PR 01-NOV-2000; 2000US-245084P.  
XX  
PA (AVAL-) AVALON PHARM.  
XX  
PI Young PE, Augustus M, Carter KC, Ebner R, Endress G, Horrigan S;  
PI Soppet DR, Weaver Z;  
XX  
DR WPI; 2002-188264/24.  
XX  
XX  
PT Screening for anti-neoplastic agent involves exposing cells to a  
PT chemical agent to be tested for anti-neoplastic activity, and  
PT determining a change in expression of a gene of a signature gene set -  
XX  
PS Claim 1; SEQ ID 2299; 44pp; English.  
XX  
CC The present invention describes a method (M1) for screening for an  
CC anti-neoplastic agent. The method involves exposing cells to a chemical  
CC agent to be tested for anti-neoplastic activity, determining a change in  
CC expression of at least one gene (I) of a signature gene set, where (I)  
CC comprises a sequence (S) selected from 8447 sequences (given in ABU61664  
CC to ABU70110), or is at least 95% identical to (S), where a change in  
CC expression is indicative of anti-neoplastic activity. (I) has cytostatic  
CC activity and can be used in gene therapy. M1 can be used for screening  
CC an anti-neoplastic agent, and can be used for producing a product which  
CC is the data collected with respect to the anti-neoplastic agent as a  
CC result of M1, and the data is sufficient to convey the chemical  
CC structure and/or properties of the agent. M1 can be used in the  
CC treatment of cancer such as colon, breast, stomach, lung, thyroid,  
CC esophageal, ovarian, kidney, prostate or pancreatic cancer,  
CC adenocarcinoma, carcinoma, clear cell cancer, infiltrating ductal cancer,  
CC infiltrating lobular cancer, squamous cell carcinoma, neuroendocrine  
CC carcinoma, papillary carcinoma and Wilm's tumour.  
XX  
SQ Sequence 306 BP; 98 A; 55 C; 40 G; 113 T; 0 other;  
XX  
Query Match 8.0%; Score 40; DB 24; Length 306;  
Best Local Similarity 59.8%; Pred. No. 0.19;  
Matches 67; Conservative 0; Mismatches 45; Indels 0; Gaps 0;  
XX  
QY 119 TTTTCTTTTACAGGCATATGAAATAAAACATATTTTGCAGCATTTGACACTTTTTC 178  
|||||  
119 TTTTCTTTTACAGGCATATGAAATAAAACATATTTTGCAGCATTTGACACTTTTTC 178  
|||||

Db 1 TTTTCTTTTACAGGCATATGAAATAAAACATATTTTGCAGCATTTGACACTTTTTC 60  
QY 179 CTTTCTAGAAATTTCTAAACCTCTGCATTTGGTGGAGACATTTGAGTACATTT 230  
|||||  
Db 61 CATCTTTTAAACAGTCTACACCGAAACATTTTGGAAACATCTTTTCCCTTT 112  
|||||

RESULT 10  
ABL33688  
ID ABL33688 standard; DNA; 9646 BP.  
XX AC ABL33688;  
XX  
DT 26-MAR-2002 (first entry)  
XX  
DE Human immune system associated gene SEQ ID NO: 1661.  
XX  
KW Human; immune system disease; cytosine methylation; antiasthmatic;  
KW antiarteriosclerotic; antianaemic; cytosolic; neurotropic;  
KW neuroprotective; anti-HIV; anticonvulsant; ophthalmological;  
KW antirheumatic; antiarthritic; antidiabetic; antipsoriatic;  
KW antiinflammatory; cancer; eye disease; arteriosclerosis; anaemia;  
KW acute myeloid leukaemia; Alzheimer's disease; AIDS; epilepsy;  
KW neurofibromatosis; rheumatoid arthritis; psoriasis; bowel disease;  
KW gene; ds.  
XX  
OS Homo sapiens.  
XX  
PN WO200200928-A2.  
XX  
PD 03-JAN-2002.  
XX  
XX 02-JUL-2001; 2001WO-EP07537.  
XX  
PR 30-JUN-2000; 2000DE-1032529.  
PR  
XX 01-SEP-2000; 2000DE-1043826.  
XX  
PA (EPIG-) EPIGENOMICS AG.  
XX  
XX Olek A, Piepenbrock C, Berlin K;  
XX  
DR WPI; 2002-130909/17.  
XX  
XX Nucleic acid comprising fragment of chemically modified gene, useful  
PT for diagnosis and treatment of diseases associated with abnormal  
PT cytosine methylation -  
PS  
XX Claim 1; SEQ ID NO 1661; 32pp + Sequence Listing; German.  
XX  
CC The present invention provides a number of human immune system associated  
CC genes which are modified by the methylation of cytosines. The sequences  
CC can be used in the diagnosis and treatment of immune system disorders,  
CC including eye diseases such as retinopathy, neovascular glaucoma and  
CC macular degeneration, arteriosclerosis, anaemia, cancer, acute myeloid  
CC leukaemia, Alzheimer's disease, AIDS, epilepsy, neurofibromatosis,  
CC rheumatoid arthritis, psoriasis and inflammatory/ulcerative bowel  
CC diseases. The present sequence is a gene of the invention.  
XX  
SQ Sequence 9646 BP; 2765 A; 150 C; 2136 G; 4595 T; 0 other;  
XX  
Query Match 7.9%; Score 39.8; DB 24; Length 9646;  
Best Local Similarity 57.7%; Pred. No. 0.95;  
Matches 71; Conservative 0; Mismatches 52; Indels 0; Gaps 0;  
XX  
QY 72 AGAATTCCTGCTGCTACTTTTCTCCAGGAGCACTCTCTGATATTTTATTTTACA 131  
|||||  
Db 6728 AATTTGTTGTTGATATTTTATTTTATTTAGGTTTATTTAGTTTATTTTGGG 6787  
|||||

QY 132 GGCATATGAATAAAACATATTTTGCAGCATTTGACACTTTTCTTTCTAGAAAT 191  
|||||  
Db 6788 GTGATTTTATTATAATGTAATTTGATTGATAGGAATTTATTTTATTTTAGTAGT 6847  
|||||  
QY 192 TCT 194

```
Db      6848 TTT 6850

RESULT 11
AAC66096
ID AAC66096 standard; cDNA; 2360 BP.
XX
AC AAC66096;
XX
XX
DT 13-FEB-2001 (first entry)
XX
DE
KW Cold resistance; transgenic plant; sucrose decomposition; rice;
KW Cold resistance; transgenic plant; sucrose decomposition; rice;
XX Oryza sativa.
XX OS
XX JP2000245279-A.
XX PN
XX 12-SEP-2000.
XX PD
XX
XX pSS3 cDNA sequence used in cold resistant plant production SEQ ID 6.
XX
XX Cold resistance; transgenic plant; sucrose decomposition; rice; ss.
XX
XX Oryza sativa.
XX OS
XX JP2000245279-A.
XX PN
XX 12-SEP-2000.
XX PD
XX
XX 01-MAR-1999; 99JP-0052102.
XX PF
XX 01-MAR-1999; 99JP-0052102.
XX PR
XX (MITA ) MITSUI CHEM INC.
XX PA
XX WPI; 2000-675173/66.
XX DR
XX P-PSDB; AAY85664.
XX DR
XX
PT Novel method for the reinforcement of cold resistance in a plant
PT comprising introducing a vector encoding an enzyme that decomposes
PT sucrose into the plant -
XX
XX Example 1; Page 15-18; 22pp; Japanese.
XX
XX This invention relates to a method for the reinforcement of cold
XX resistance in a plant, comprising introducing an expression vector having
XX a DNA encoding an enzyme for decomposing sucrose connected downstream to
XX a promoter expressible in the plant, and expressing the enzyme in the
XX plant body. Included in the invention are an expression vector used in
XX the method; a transformed plant carrying the expression vector; and a
XX transformed rice carrying the above expression vector. The method is used
XX for reinforcing cold resistance in a plant. The present sequence
XX represents cDNA used in the production of cold resistant plants by the
XX method of the invention.
XX
XX Sequence 2360 BP; 595 A; 556 C; 599 G; 610 T; 0 other;
XX
Query Match 7.9%; Score 39.6; DB 21; Length 2360;
Best Local Similarity 54.0%; Pred. No. 0.59;
Matches 81; Conservative 0; Mismatches 69; Indels 0; Gaps 0;
Qy 173 TTTTTCCTTTTCTAGAAAATCTAAACCTCTGACATTTGTTGGAGACATTGAGTACATTTT 232
Db 742 TTTGCCAATCCCATGTGTTGGGATACCTGATCTGTTGTCAGTTGTGTACATTTG 801
Qy 233 TCCCATATCCCTACTTTTCAGAAAGATTTCCTGCTCGTTTCACTTAACATTTGCTGATGC 292
Db 802 GACCAAGTCCGCGCTTTGGAGAATGAGATGCTTTTGAGGATCAAGCAGACGAGCCTTGAT 861
Qy 293 GTGAGTCTTTTCTCCCTCATCTCTTTTCAGG 322
Db 862 ATCACACTAAGATCCCTCATTTGTAACACAGG 891

RESULT 12
AAC66090
ID AAC66090 standard; cDNA; 2494 BP.
XX
AC AAC66090;
XX
XX
DT 13-FEB-2001 (first entry)
XX
DE
KW Human immune/haematopoietic antigen genomic sequence SEQ ID NO:24714.
KW Human; immune; haematopoietic; immune/haematopoietic antigen; cancer;
XX cytostatic; gene therapy; vaccine; metastasis; ds.
XX
XX Homo sapiens.
XX OS
XX W0200157182-A2.
XX PN

13-FEB-2001 (first entry)
Rice sucrose synthase cDNA sequence.
Cold resistance; transgenic plant; sucrose decomposition; rice;
sucrose synthase; ss.
Oryza sativa.
JP2000245279-A.
12-SEP-2000.
01-MAR-1999; 99JP-0052102.
01-MAR-1999; 99JP-0052102.
(MITA ) MITSUI CHEM INC.
WPI; 2000-675173/66.
P-PSDB; AAY85664.
Novel method for the reinforcement of cold resistance in a plant
comprising introducing a vector encoding an enzyme that decomposes
sucrose into the plant -
Claim 1; Page 8-11; 22pp; Japanese.
This invention relates to a method for the reinforcement of cold
resistance in a plant, comprising introducing an expression vector having
a DNA encoding an enzyme for decomposing sucrose connected downstream to
a promoter expressible in the plant, and expressing the enzyme in the
plant body. Included in the invention are an expression vector used in
the method; a transformed plant carrying the expression vector; and a
transformed rice carrying the above expression vector. The method is used
for reinforcing cold resistance in a plant. The present sequence
represents cDNA encoding sucrose synthase used in the method.
Sequence 2494 BP; 622 A; 599 C; 628 G; 645 T; 0 other;
Query Match 7.9%; Score 39.6; DB 21; Length 2494;
Best Local Similarity 54.0%; Pred. No. 0.61;
Matches 81; Conservative 0; Mismatches 69; Indels 0; Gaps 0;
Qy 173 TTTTTCCTTTTCTAGAAAATCTAAACCTCTGACATTTGTTGGAGACATTGAGTACATTTT 232
Db 876 TTTGCCAATCCCATGTGTTGGGATACCTGATCTGTTGTCAGTTGTGTACATTTG 935
Qy 233 TCCCATATCCCTACTTTTCAGAAAGATTTCCTGCTCGTTTCACTTAACATTTGCTGATGC 292
Db 936 GACCAAGTCCGCGCTTTGGAGAATGAGATGCTTTTGAGGATCAAGCAGACGAGCCTTGAT 995
Qy 293 GTGAGTCTTTTCTCCCTCATCTCTTTTCAGG 322
Db 996 ATCACACTAAGATCCCTCATTTGTAACACAGG 1025

RESULT 13
AAC69902/c
ID AAC69902 standard; DNA; 648 BP.
XX
XX AAC69902;
XX
XX 06-NOV-2001 (first entry)
XX
XX Human immune/haematopoietic antigen genomic sequence SEQ ID NO:24714.
XX Human; immune; haematopoietic; immune/haematopoietic antigen; cancer;
XX cytostatic; gene therapy; vaccine; metastasis; ds.
XX
XX Homo sapiens.
XX OS
XX W0200157182-A2.
XX PN
```

XX 09-AUG-2001.  
XX 17-JAN-2001; 2001WO-US01354.  
XX 31-JAN-2000; 2000US-0179065.  
XX 04-FEB-2000; 2000US-0180628.  
XX 24-FEB-2000; 2000US-0184664.  
XX 02-MAR-2000; 2000US-0186350.  
XX 16-MAR-2000; 2000US-0189874.  
XX 17-MAR-2000; 2000US-0190076.  
XX 18-APR-2000; 2000US-0198123.  
XX 19-MAY-2000; 2000US-0205515.  
XX 07-JUN-2000; 2000US-0209467.  
XX 28-JUN-2000; 2000US-0214886.  
XX 30-JUN-2000; 2000US-0215135.  
XX 07-JUL-2000; 2000US-0216647.  
XX 07-JUL-2000; 2000US-0216880.  
XX 11-JUL-2000; 2000US-0217487.  
XX 11-JUL-2000; 2000US-0217496.  
XX 14-JUL-2000; 2000US-0218290.  
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XX 17-NOV-2000; 2000US-0249300.  
XX 01-DEC-2000; 2000US-0250160.  
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XX 05-DEC-2000; 2000US-0251030.  
XX 05-DEC-2000; 2000US-0251988.  
XX 05-DEC-2000; 2000US-0256719.  
XX 06-DEC-2000; 2000US-0251479.  
XX 08-DEC-2000; 2000US-0251856.  
XX 08-DEC-2000; 2000US-0251868.  
XX 08-DEC-2000; 2000US-0251869.  
XX 08-DEC-2000; 2000US-0251989.  
XX 11-DEC-2000; 2000US-0251990.  
XX 05-JAN-2001; 2001US-02559678.  
(HUMA-) HUMAN GENOME SCI INC.  
Rosen CA, Barash SC, Ruben SM;  
XX  
XX  
XX  
XX



```
DR WPI; 2001-483426/52.
XX
XX Nucleic acids encoding human immune/hematopoietic antigen polypeptides,
PT useful for preventing, diagnosing and/or treating cancers and
PT metastasis -
XX
XX Disclosure; SEQ ID NO 24714; 3071pp + Sequence Listing; English.
XX
XX AAK54951 to AAK64702 encode the human immune/hematopoietic antigen (I)
CC amino acid sequences given in AAK82170 to AAK91921. (I) have cytostatic
CC activity, and can be used in gene therapy and vaccine production. (I)
CC proteins and polynucleotides may be used in the prevention, diagnosis and
CC treatment of diseases associated with inappropriate (I) expression. For
CC example, they may be used to treat disorders associated with decreased
CC expression by rectifying mutations or deletions in a patient's genome
CC that affect the activity of (I) by expressing inactive proteins or to
CC supplement the patient's own production of (I). Additionally, (I)
CC polynucleotides may be used to produce the secreted (I), by inserting
CC the nucleic acids into a host cell and culturing the cell to express the
CC protein. (I) proteins and polynucleotides may be used to prevent,
CC diagnose and treat immune/hematopoietic-related diseases, especially
CC cancers and cancer metastases of hematopoietic-derived cells. AAK64703
CC to AAK87694 represent human immune/hematopoietic antigen genomic
CC sequences from the present invention. AAK54942 to AAK54950 and AAK82169
CC represent sequences used in the exemplification of the present invention.
XX
XX Sequence 648 BP; 233 A; 96 C; 122 G; 197 T; 0 other;
SQ
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Best Local Similarity 53.3%; Pred. No. 0.66;
Matches 105; Conservative 0; Mismatches 89; Indels 3; Gaps 1;
QY 86 TACTTTTCTCCAGAGAACCTTCCTTGATATTTTTCCTTTCTAGAAATCTTAACCTCTGAC 145
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282 TATATTTGATGTCAGCAACCTTCCTTGATATTTTTCCTTTCTACTATATTAAGCCCACT 223
QY 146 AACTATATTTGCAGCATGTACACTTTTTCCTTTCTAGAAATCTTAACCTCTGAC 205
Db ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
222 ACTTACTCTGTGTGTTGTTCTTTTTCCTTTCTACTATATTAAGCCCACT 163
QY 206 ATTG--GTGAGACATTTGAGTACATTTTTCCTATCCCTACTTTTCAGAGGATTTT 262
Db ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
162 TGAGAAAGAGATAATAGTAATATTTTGTGATTTGTAAGTCTTCAGAGCATACA 103
QY 263 CTCGCTCGTTCACHTA 279
Db ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
102 GACTCTCGATGACGTA 86
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ID AAV54123 standard; DNA; 2857 BP.
XX
XX AAV54123;
AC
XX
XX 17-DEC-1998 (first entry)
DT
XX
DE Rabbit membrane protein BA0306 coding sequence.
XX
XX Membrane protein; BA0306; BA2303; arteriosclerosis; coronary restenosis;
KW therapy; rabbit; ds.
XX
XX Oryctolagus sp.
OS
XX
XX Key Location/Qualifiers
FH 415..2199
CDS /*tag= a
FT /transl_except= (pos: 745..747, aa: Xaa)
FT /note= "Xaa= unspecified amino acid"
XX
XX WO9838305-A1.
FN
XX 03-SEP-1998.
PD
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XX
XX 27-FEB-1998; 98WO-JP00835.
PF
XX
XX 25-FEB-1998; 98JP-0062263.
PR
XX 28-FEB-1997; 97JP-0062259.
PR
XX (NISR) JAPAN TOBACCO INC.
PA
XX
XX Nakamura Y, Tanaka T, Tsukada S;
PI
XX
XX WPI; 1998-481206/41.
DR
XX P-PSDB; AAW74579.
XX
XX Membrane protein(s) BA0306 and BA2303 - useful for, e.g. treatment
PT and prevention of arteriosclerosis and restenosis
PT
XX
XX Example 4; Page 80-85; 141pp; Japanese.
PS
XX
XX This sequence represents the DNA encoding the rabbit
CC membrane protein BA0306. This sequence was used to isolate the human
CC membrane proteins BA0306 and BA2303 of the invention. The two membrane
CC proteins are specifically expressed in mammals during arteriosclerosis
CC and coronary restenosis. The membrane proteins, fragments of them, and
CC antibodies against them are useful in the treatment and prevention of
CC arteriosclerosis and restenosis. Transgenic mice expressing the
CC extracellular region of the membrane proteins are useful as models for
CC studying these disorders.
XX
XX Sequence 2857 BP; 787 A; 553 C; 588 G; 927 T; 2 other;
SQ
Query Match 7.6%; Score 38.2; DB 19; Length 2857;
Best Local Similarity 52.9%; Pred. No. 1.6;
Matches 82; Conservative 0; Mismatches 73; Indels 0; Gaps 0;
QY 37 GAAGGCGATTGGGAGTCAGCTCCTAAGTAAGCTCCAGAAATCTCTGCTACTTTTCCCT 96
Db ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
2451 GAATGGACACAATGATGATCTTAATGAGACCAAAAGTACTTCTGTGTTTCCCTTCG 2510
QY 97 CCAGGAAGCAACTTCCTTTGATATTTTTCCTTTTACAGGCATATGAATAAACTATATTT 156
Db ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
2511 TCAGAAAGCATCTCCATTGTAATATGATTTACATGTTTATTACAAAGATCCAATGAA 2570
QY 157 GCAGCATTTGTACACTTTTTCCTTTTCTAGAAAT 191
Db ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
2571 AAATTTTGTCCATTTTTCATAGCCCTAAAGAT 2605
RESULT 15
AAS45388
ID AAS45388 standard; DNA; 15732 BP.
XX
XX AAS45388;
AC
XX
XX 18-DEC-2001 (first entry)
DT
XX
DE Chemically pretreated genomic DNA associated with cell cycle #47.
XX
XX Cell cycle; human; CpG dinucleotide; cytosine methylation; HIV; aging;
KW human immunodeficiency virus; neurodegenerative disorder; solid tumour;
KW graft-versus-host disease; glomerular disease; Lewy body disease; cancer;
KW arthritis; arteriosclerosis; anti-HIV; neuroprotective; antiarthritic;
KW immunosuppressive; antitumour; cytostatic; antiarteriosclerotic; ds;
KW PCR primer.
XX
XX Homo sapiens.
OS
XX
XX WO200168911-A2.
FN
XX
XX 20-SEP-2001.
PD
XX
XX 15-MAR-2001; 2001WO-EP02945.
PF
XX
XX 15-MAR-2000; 2000DE-1013847.
PR
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PR 06-APR-2000; 2000DE-1019058.
PR 07-APR-2000; 2000DE-1019173.
PR 30-JUN-2000; 2000DE-1032529.
PR 01-SEP-2000; 2000DE-1043826.
XX
XX (EPiG-) EPIGENOMICS AG.
XX
XX PI Olek A, Piepenbrock C, Berlin K;
XX
XX DR WPI; 2001-602751/68.
XX
XX Designing primers and probes for analysing diseases associated with
XX cytosine methylation state e.g. arthritis, cancer, aging,
XX PT arteriosclerosis comprising fragments of chemically modified genes
XX PT associated with cell cycle -
XX
XX Claim 1; SEQ ID No 93; 28pp; English.
XX
XX Sequences AAS45296-AAS45520 represent chemically pretreated genomic DNA
XX molecules associated with the cell cycle and specific PCR primers of the
XX invention. The sequences are useful for detecting the methylation state
XX of all CpG dinucleotides in a sequence and therefore for analysing
XX associated diseases. By analysing cytosine methylations in the pretreated
XX DNA, genetic and/or epigenetic parameters for the diagnosis and therapy
XX of existing diseases or the predisposition to specific diseases can be
XX ascertained. The parameters may be compared to another set of genetic
XX and/or epigenetic parameters, the differences serving as basis for
XX diagnosis and/or prognosis events which are disadvantageous to patients.
XX The sequences of the invention are useful for the diagnosis and therapy
XX of HIV infection, neurodegenerative disorders, graft-versus-host disease,
XX aging, glomerular disease, Lewy body disease, arthritis,
XX arteriosclerosis, solid tumours and cancers.
XX
SQ Sequence 15732 BP; 4638 A; 70 C; 2672 G; 8352 T; 0 other;

Query Match          7.6%; Score 38.2; DB 22; Length 15732;
Best Local Similarity 58.3%; Pred. No. 3.3;
Matches 67; Conservative 0; Mismatches 48; Indels 0; Gaps 0;

QY 119 TTTTCTTTTTCAGGCATATGAATAAAACTATATTTTGCAGCATTTGACACTTTTTC 178
Db      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
10649 TTTTATTTTATTATTATGATAAAAAAATATATTTTGTATGATTTTAAACGTTTATA 10708

QY 179 CTTTCTAGAAATTCCTAACCTCTGACATTTGTCAGACATTTGACATATTTTTC 233
Db      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
10709 TTTATTAATATTGTGGTATTTTAAATTGCGTATATGTTGGGATAATGTTT 10763
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Search completed: November 7, 2003, 06:15:41  
Job time : 157.728 secs

GenCore version 5.1.6  
Copyright (c) 1993 - 2003 Compugen Ltd.

OM nucleic - nucleic search, using sw model

Run on: November 6, 2003, 21:13:51; Search time 155.728 Seconds  
(without alignments)  
8684.478 Million cell updates/sec

Title: US-09-939-209A-3\_COPY\_15000\_15500

Perfect score: 501

Sequence: 1 tggcagagaactctctgga.....tcaacctgatggagaaggat 501

Scoring table: IDENTITY\_NUC

Gapop 10.0, Gapext 1.0

Searched: 2552756 seqs, 1349719017 residues

Total number of hits satisfying chosen parameters: 5105512

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-Processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database: N Geneseq 19Jun03.\*

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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

#### SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	501	100.0	20300	24	ABK47337 Genomic nucleotide
2	113	22.6	800	25	ACA56493 Human signalling p
3	113	22.6	1238	22	AAL26766 Human breast cancer
4	113	22.6	2934	24	ABK47336 cDNA encoding huma
5	113	22.6	3165	25	ABX63113 Human cDNA #113 di
6	88.2	17.6	201	18	AAU50916 Rat brain xgse-1 g
7	60	12.0	60	24	ABNA2053 Human spliced tran
8	52.6	10.5	10427	24	ABK69943 Human secreted pro

9	49	9.8	378	24	ABL99976	Rat disease associ
10	49	9.8	543	22	ABL99975	Rat disease associ
11	49	9.8	618	22	ABA09731	Human bone marrow
12	49	9.8	817	24	ABL99979	Rat disease associ
13	48	9.6	540	24	ABA92505	Human regulator of
14	48	9.6	606	21	AAA39678	Cytokine response
15	48	9.6	746	19	AAV34780	Human RATH1.1 DNA.
16	48	9.6	2383	17	AAT11418	p53 response prote
17	48	9.6	2406	18	AA743376	Human cytokine res
18	48	9.6	2406	21	AAA39660	Human CR1 cDNA. H
19	48	9.6	2434	24	AA94857	Human DNA sequence
20	47.4	9.5	745	19	AAV38084	Human regulator of
21	47.4	9.5	1691	21	AAZ36910	cDNA encoding a re
22	47.4	9.5	1923	20	AAZ51745	DNA encoding a hum
23	47.4	9.5	1923	24	AAQ92597	Human secreted pro
24	47.4	9.5	2075	25	ABX74396	Human cDNA sequenc
25	46.4	9.3	597	22	AAZ57422	Human RGS8LIKE pol
26	46.4	9.3	939	24	ABZ11471	Human polynucleoti
27	45.8	9.1	955	22	AAI97781	Human neuroblastom
28	44.8	8.9	2272	19	AAV34779	Mouse RATH1.1 DNA.
29	43.8	8.7	1396	23	ABV21629	Human prostate exp
30	43.8	8.7	1396	23	ABV27449	Human prostate exp
31	43.8	8.7	7345	24	ABL62354	Colon adenocarcino
32	43.2	8.6	1978	21	AAZ79846	Human secreted pro
33	42.2	8.4	411	23	ABX35169	Bovine EST associa
34	41.8	8.3	1345	24	ABT10881	Human breast cancer
35	41.8	8.3	1345	24	ABK93834	Human cDNA differe
36	41.8	8.3	1345	25	ACC46750	Human COPD related
37	41.8	8.3	1345	25	ACA56680	Signalling pathway
38	41.8	8.3	1364	21	AAFI6132	Human prostate can
39	41.6	8.3	2638	22	AAH02909	Human shear stress
40	41.6	8.3	2638	24	ABL62703	Colon adenocarcino
41	41.6	8.3	2638	24	ABL66645	Lung cancer relate
42	41.6	8.3	2638	25	ACA56492	Human signalling p
43	41.6	8.3	2874	25	ABX72245	Human NOVX polynuc
44	41	8.2	342	22	AAK54581	Human haematologic
45	41	8.2	342	22	AAK54733	Human haematologic

#### ALIGNMENTS

##### RESULT 1

ABK47337  
ID ABK47337 standard; DNA; 20300 BP.

XX AC ABK47337;

XX DT 18-JUN-2002 (first entry)

XX DE Genomic nucleotide sequence encoding human RGS-4 protein.

XX KW RGS-4; schizophrenia; human; regulator of G protein signalling 4;  
neuroleptic; polymorphism; schizotypy; schizoid; chromosome 1q21-22;  
gene; ds.

XX OS Homo sapiens.

XX FH Key

FT variation

FT Location/Qualifiers

FT replace (4121,T)

FT /\*tag= a

FT /standard\_name= "Single-nucleotide polymorphism"

FT replace (4123,A)

FT /\*tag= b

FT /standard\_name= "Single-nucleotide polymorphism"

FT replace (4368,C)

FT /\*tag= c

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FT replace (4621,C)

FT /\*tag= d

FT /standard\_name= "Single-nucleotide polymorphism"

FT replace (4790,T)

FT /\*tag= e

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FN W0200216653-A2.  
XX  
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XX  
XX 24-AUG-2001; 2001WO-US266622.  
XX  
XX 24-AUG-2000; 2000US-228021P.  
XX
```

```
PA (UYPI-) UNIV PITTSBURGH.  
XX  
XX Levitt PR, Mirnics K, Kodavali VC, Nimgaonkar VL;  
XX  
XX WPI; 2002-292070/33.  
XX  
XX Diagnosing, assessing susceptibility and treating schizophrenia,  
XX involves observing regulator of G-protein signalling 4, RGS4 levels in a  
XX subject -  
XX  
XX Claim 1; Page 20-33; 112pp; English.  
XX  
XX This invention relates to a novel method for diagnosing schizophrenia  
XX or determining susceptibility to schizophrenia in a human. The method  
XX comprises obtaining from a patient a DNA sample and detecting variations  
XX in the regulator of G-protein signalling 4 (RGS4) gene. Alternatively,  
XX the method involves measuring RGS4 mRNA or protein levels in a tissue  
XX sample from the patient and determining if there is a reduced level.  
XX The method of the invention is useful for diagnosing and determining  
XX susceptibility to schizophrenia. The invention also comprises a method  
XX that is useful for treating schizophrenia which includes a prophylactic  
XX treatment. The method of genotyping polymorphic variants in the RGS-4  
XX gene is applied to diagnosing pathologies of the schizophrenic spectrum,  
XX such as in particular schizotypy, schizoid individuals, etc. This  
XX method offers the possibility of diagnosing schizophrenia by a  
XX biological test and no longer exclusively by clinical evaluations.  
XX The present sequence represents the genomic DNA encoding the human  
XX regulator of G-protein signalling 4 (RGS4) protein used in the method of  
XX the invention. The gene for the RGS4 protein is located on human  
XX chromosome 1q21-22.  
XX  
XX Sequence 20300 BP; 6157 A; 4102 C; 3775 G; 6266 T; 0 other;  
XX  
XX Query Match 100.0%; Score 501; DB 24; Length 20300;  
XX Best Local Similarity 100.0%; Pred. No. 1.5e-146;  
XX Matches 501; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
Qy 1 TGGCAGAGAACTCTCTGGATCTTAGTGAAGGTTCTTAGAATAGTGGAGCTGACTATCATATA 60  
Db 15000 TGGCAGAGAACTCTCTGGATCTTAGTGAAGGTTCTTAGAATAGTGGAGCTGACTATCATATA 15059  
Qy 61 ATCTTGACACCCCAATAAATCAGTTTTTAAAAAATCTTTTATCCATGGGCTTAC 120  
Db 15060 ATCTTGACACCCCAATAAATCAGTTTTTAAAAAATCTTTTATCCATGGGCTTAC 15119  
Qy 121 CATAACCTCCCTGCATGAATTTTTCGATGAATCTCCCAATTTGTTAGACAGACAGAA 180  
Db 15120 CATAACCTCCCTGCATGAATTTTTCGATGAATCTCCCAATTTGTTAGACAGACAGAA 15179  
Qy 181 GATCTTGGCCCTGCTCTCTCTAAAGCAGAAAGGTTCAATCTGAACCTTTTCACTCTCA 240  
Db 15180 GATCTTGGCCCTGCTCTCTCTAAAGCAGAAAGGTTCAATCTGAACCTTTTCACTCTCA 15239  
Qy 241 CATGTGCGAAGGAGGCCCAATGTCACTTTTCTTTTCTTTTCTGAAATACAGAGGGTG 300  
Db 15240 CATGTGCGAAGGAGGCCCAATGTCACTTTTCTTTTCTTTTCTGAAATACAGAGGGTG 15299  
Qy 301 CACTGCCACTTACAAGTCACTACAAGCATACAGGCTTGATCTCTCAACAGGATATAGG 360  
Db 15300 CACTGCCACTTACAAGTCACTACAAGCATACAGGCTTGATCTCTCAACAGGATATAGG 15359  
Qy 361 TCTAATGAAGCCTTGGCCCTTGGCCCTCAGGTGAACCTGATTTCTTGACACAGGGAAGAG 420  
Db 15360 TCTAATGAAGCCTTGGCCCTTGGCCCTCAGGTGAACCTGATTTCTTGACACAGGGAAGAG 15419  
Qy 421 ACAAGCGGAACATGCTAGAGCCTCAATAAATCCTGCTTTTGTAGTCCGCCACAGAGAGATT 480  
Db 15420 ACAAGCGGAACATGCTAGAGCCTCAATAAATCCTGCTTTTGTAGTCCGCCACAGAGAGATT 15479  
Qy 481 TTCAACCTGATGGAGAAGGAT 501  
Db 15480 TTCAACCTGATGGAGAAGGAT 15500
```

RESULT 2  
ACA56493  
ID ACA56493 standard; cDNA; 800 BP.  
XX  
AC ACA56493;  
XX  
DT 06-JUN-2003 (first entry)  
XX  
DE Human signalling pathway polynucleotide probe SEQ ID NO 1091.  
XX  
KW Human; probe; ss; array element; Parkinson's disease;  
KW signalling pathway population; cancer; adenocarcinoma; leukaemia;  
KW immunopathy; AIDS; asthma; neuropathy; Alzheimer's disease; microarray.  
XX  
OS Homo sapiens.  
XX  
FN US6500938-B1.  
XX  
PD 31-DEC-2002.  
XX  
PF 30-JAN-1998; 98US-0016434.  
XX  
PR 30-JAN-1998; 98US-0016434.  
XX  
PA (INCY-) INCYTE GENOMICS INC.  
XX  
PI Au-Young J, Seilhamer JJ;  
XX  
DR WPI; 2003-352189/33.  
XX  
PT Combination of polynucleotide probes, useful as array elements in a  
PT microarray for monitoring the expression of a number of target  
PT polynucleotides -  
XX  
PS Claim 1; SEQ ID NO 1091; 65pp; English.  
XX  
CC The invention relates to a combination which, comprises a number of  
CC polynucleotide probes comprising a sequence selected from one of the 1490  
CC sequences mentioned in the specification. The combination is useful as an  
CC array element in a microarray for monitoring the expression of a number  
CC of target polynucleotides. The microarray is particularly useful in the  
CC diagnosis and treatment of cancer and immunopathology and neuropathology.  
CC The microarray is useful in diagnostics and treatment regimens, drug  
CC discovery and development, toxicological and carcinogenicity studies,  
CC forensics and pharmacogenomics. The microarray is also useful for  
CC monitoring progression of diseases and for developing sophisticated  
CC profiles for the effects of currently available therapeutic drugs. The  
CC combination is also useful for purifying a subpopulation of mRNAs, cDNAs  
CC and genomic fragments and in research and diagnostic applications. The  
CC array can detect changes in expression in a large number of genes coding  
CC for different signalling pathway populations which can be used to diagnose  
CC various diseases including cancer e.g. adenocarcinoma and leukaemia,  
CC immunopathies e.g. AIDS and asthma, neuropathies e.g. Alzheimer's disease  
CC and Parkinson's disease. The present sequence represents a polynucleotide  
CC probe of the invention.  
CC Note: The sequence data for this patent did not form part of the printed  
CC specification but was obtained in electronic format directly from USPTO  
CC at [seqdata.uspto.gov/sequence.html?docID=06500938B1](http://seqdata.uspto.gov/sequence.html?docID=06500938B1).  
XX  
SQ Sequence 800 BP; 241 A; 181 C; 195 G; 183 T; 0 other;  
Query Match 22.6%; Score 113; DB 25; Length 800;  
Best Local Similarity 100.0%; Pred. No. 4.3e-25;  
Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 389 AGGTGAACCTGGATTCTTGACCGAGAGACAAGCCGAGACATGCTAGAGCTACAA 448  
Db 474 AGGTGAACCTGGATTCTTGACCGAGAGACAAGCCGAGACATGCTAGAGCTACAA 533  
QY 449 TAACCTGCTTTGATGAGGCCCGAGAGAGATTTTCAACCTGATGGAGAGGAT 501  
Db 534 TAACCTGCTTTGATGAGGCCCGAGAGAGATTTTCAACCTGATGGAGAGGAT 586

RESULT 3  
AAL26766  
ID AAL26766 standard; cDNA; 1238 BP.  
XX  
AC AAL26766;  
XX  
DT 07-DEC-2001 (first entry)  
XX  
DE Human breast cancer expressed polynucleotide 19223.  
XX  
KW Human; breast cancer; cell marker; cytostatic; ss.  
XX  
OS Homo sapiens.  
XX  
FN WO2000151628-A2.  
XX  
PD 19-JUL-2001.  
XX  
PF 10-JAN-2001; 2001WO-US00798.  
XX  
PR 14-JAN-2000; 2000US-0176077.  
XX  
PR 14-MAR-2000; 2000US-0189167.  
XX  
PR 24-MAR-2000; 2000US-0192099.  
XX  
PR 29-MAR-2000; 2000US-0193480.  
XX  
PR 15-MAY-2000; 2000US-0205230.  
XX  
PR 09-JUN-2000; 2000US-0211315.  
XX  
PR 25-JUL-2000; 2000US-0220534.  
XX  
PA (MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.  
XX  
PI Lillie J, Xu Y, Wang Y, Steinmann K;  
XX  
DR WPI; 2001-451856/48.  
XX  
PT New peptide useful as a marker for the diagnosis of breast cancer -  
XX  
PS Claim 1; Page 3683-3684; 3695pp; English.  
XX  
CC The invention relates to human breast cancer expressed polynucleotides  
CC (AAL07544-AAL26789) and methods of assessing whether a patient is  
CC afflicted with breast cancer by examining the correlation between the  
CC expression of certain markers and the cancerous state of breast cells.  
CC The polynucleotides and encoded polypeptides are potential markers for  
CC detecting, diagnosing, monitoring, characterizing treating and  
CC potentially preventing breast cancer. The polynucleotides and encoded  
CC polypeptides are also useful for isolating compounds with cytostatic  
CC activity.  
XX  
SQ Sequence 1238 BP; 386 A; 265 C; 307 G; 274 T; 6 other;  
Query Match 22.6%; Score 113; DB 22; Length 1238;  
Best Local Similarity 100.0%; Pred. No. 5.3e-25;  
Matches 113; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 389 AGGTGAACCTGGATTCTTGACCGAGAGACAAGCCGAGACATGCTAGAGCTACAA 448  
Db 674 AGGTGAACCTGGATTCTTGACCGAGAGACAAGCCGAGACATGCTAGAGCTACAA 733  
QY 449 TAACCTGCTTTGATGAGGCCCGAGAGAGATTTTCAACCTGATGGAGAGGAT 501  
Db 734 TAACCTGCTTTGATGAGGCCCGAGAGAGATTTTCAACCTGATGGAGAGGAT 786  
RESULT 4  
ABK47336  
ID ABK47336 standard; DNA; 2934 BP.  
XX  
AC ABK47336;  
XX  
DT 18-JUN-2002 (first entry)  
XX



Db	679	AGGTGAACCTGGATTCTTCGACCGGGGAGAGACAGCCGGACATCTCTAGACCTACAA	739
Qy	449	TAACTCTGCTTTGATGAGGCCAGAGAAGATTTTCAACCTGATGGAGAAGAT	501
Db	739	TAACTCTGCTTTGATGAGGCCAGAGAAGATTTTCAACCTGATGGAGAAGAT	791
RESULT 6			
AAAT50916	AAAT50916 standard; DNA; 201 BP.		
XX	AC	AAAT50916;	
XX	31-MAR-1997	(first entry)	
XX	Rat brain rgss-1 gene fragment.		
DE	XX	Regulator G-protein signalling similarity; rgss; RGS;	
KW	KW	signal transduction; transgenic animal; diagnosis; therapy;	
KW	KW	diabetes; hyperplasia; psychiatric disorder;	
KW	KW	cardiovascular disease; McCune-Albright syndrome;	
KW	KW	Albright hereditary osteopathy; sa.	
XX	OS	Rattus sp.	
XX	PN	W096384462-A1.	
XX	XX	05-DEC-1996.	
XX	XX	31-MAY-1996; 96WO-US08295.	
XX	XX	12-JAN-1996; 96US-0588258.	
PR	PR	02-JUN-1995; 95US-0460505.	
XX	XX	(MASI ) MASSACHUSETTS INST TECHNOLOGY.	
XX	PI	Horvitz HR, Koelle M;	
XX	XX	WPI; 1997-034298/03.	
DR	DR	P-PSDB; AAW10169.	
XX	XX	New isolated regulator of G-protein signalling genes - used to	
PT	PT	develop prods. for the diagnosis and treatment of G-protein related	
PT	PT	diseases and disorders e.g. diabetes, cardiovascular disease, etc	
XX	XX	Example B; Page 53; 96pp; English.	
PS	PS	Gene fragments (AAAT50916-24), designated rgss-1 to rgss-9, were	
XX	CC	isolated from rat brain cDNA using degenerate primers (AAAT50912-15)	
CC	CC	based on the conserved region of nematode EGL-10 protein (see	
CC	CC	also AAW10167 and AAW10178). EGL-10 is a member of the new RGS	
CC	CC	(Regulators of G-protein Signalling) family involved in the control	
CC	CC	of G-protein mediated effects. The rat gene fragments, designated	
CC	CC	rgss for regulator G-protein signalling similarity, encode proteins	
CC	CC	(AAW10169-77) contg. RGS conserved regions. RGS genes (see also	
CC	CC	AAAT50910-11) can be used to detect related genes, to produce RGS	
CC	CC	polypeptides in transformed host cells, and in the diagnosis and	
CC	CC	treatment of G-protein related disorders.	
XX	XX	Sequence 201 BP; 70 A; 44 C; 45 G; 41 T; 0 other;	
XX	XX	Query Match 17.6%; Score 88.2; DB 18; Length 201;	
XX	XX	Best Local Similarity 92.1%; Pred. No. 1.4e-17;	
XX	XX	Matches 93; Conservative 0; Mismatches 8; Indels 0; Gaps 0	
Qy	389	AGGTGAACCTGGATTCTTCGACCGGAGAGACAAAGCCGGAACATGCTAGAGCCTACAA	448
Db	101	AGGTGAACCTGGATTCTTCGACCGGAGAGACAAAGCCGGAACATGTTAGAGCCACGA	160
Qy	449	TAACTCTGTTGATGAGGCCAGAGAAGATTTTCAACCTG	489
Db	161	TAACTCTGTTGATGAGGCCAGAGAAGATTTTCAACCTG	201



```
RESULT 8
ABK69943
ID ABK69943 standard; DNA; 10427 BP.
XX
AC ABK69943;
XX
DT 15-JUL-2002 (first entry)
XX
DE Human secreted protein gene 69 genomic DNA fragment #2.
XX
KW Human; ds; secreted protein; gene therapy; immunosuppressive;
KW antiarthritic; antirheumatic; antiproliferative; cytostatic; cardiac;
KW vasotropic; cerebroprotective; nootropic; neuroprotective; antibacterial;
KW viricide; fungicide; ophthalmological; autoimmune disease; neoplasm;
KW rheumatoid arthritis; hyperproliferative disorder; cardiac arrest;
KW cardiovascular disorder; cerebrovascular disorder; cerebral ischaemia;
KW angiogenesis; nervous system disorder; Alzheimer's disease; infection;
KW ocular disorder; corneal infection; wound healing; skin aging;
KW epithelial cell proliferation; food additive.
XX
OS Homo sapiens.
XX
FN WO200226931-A2.
XX
PD 04-APR-2002.
XX
PF 24-SEP-2001; 2001WO-US29871.
XX
PR 25-SEP-2000; 2000US-234925P.
XX
PR 12-JAN-2001; 2001WO-US00911.
XX
PA (HUMA-) HUMAN GENOME SCI INC.
XX
PI Ruben SM, Komatsoulis G, Duan DR, Rosen CA, Moore PA, Shi Y;
PI Lafleur DW, Olsen H, Brewer LA, Florence KA, Young PE, Soppet DR;
PI Endress GA, Mucenski M, Ebner R;
XX
WPI; 2002-362489/39.
XX
Novel 71 isolated secreted polypeptides and polynucleotides encoding
the polypeptides, useful for treating Huntington's disease, sepsis,
meningitis, thrombocytopaenia, haemolytic anaemia, rheumatoid arthritis,
asthma -
XX
Example 2; Page 1452-1455; 1478pp; English.
XX
The invention relates to an isolated nucleic acid molecule (or its
fragment, homologue complement or allelic variant) encoding a human
secreted protein (and its fragment, domain, epitope, variant, secreted
form and species variant). Also included are a recombinant vector
comprising the nucleic acid, a recombinant host cell comprising the
vector, an antibody against the secreted protein, a recombinant host cell
that expresses the secreted protein and a method of identifying a binding
partner of the secreted protein. The nucleic acid and protein are used to
prevent, diagnose, treat or ameliorate a medical condition in e.g.
humans, mice, rabbits, goats, horses, cats, dogs, chickens or sheep
for example autoimmune diseases e.g. rheumatoid arthritis,
hyperproliferative disorders e.g. neoplasms of the breast or liver,
cardiovascular disorders e.g. cardiac arrest, cerebrovascular disorders
e.g. cerebral ischaemia, angiogenesis, nervous system disorders e.g.
Alzheimer's disease, infections caused by bacteria, viruses and fungi and
ocular disorders e.g. corneal infection. Many other diseases and
disorders are listed in the specification. The polypeptides can also be
used to aid wound healing in the specification. The polypeptides can also be
skin aging due to sunburn, to maintain organs before transplantation, for
supporting cell culture of primary tissues, to regenerate tissues and in
chemotaxis. The polypeptides can also be used as a food additive or
preservative to increase or decrease storage capabilities. The present
sequence represents a ds DNA fragment of the gene for a novel human
secreted protein of the invention.
XX
Sequence 10427 BP; 3337 A; 1995 C; 1829 G; 3266 T; 0 other;
SQ
```

```
Query Match
Best Local Similarity 10.5%; Score 52.6; DB 24; Length 10427;
Matches 82; Conservative 0; Mismatches 49; Indels 0; Gaps 0;
XX
QY 371 CTTGGCCCTTTGCCCTCAGGTGAACCTGGATTCTTGCACCGGGAGAGACAAGCCGGA 430
DB 5202 CCTGTTTCTTGGCCACAGGTGAATATTGACCACCTTCACTAAGGACATCAATGAAGA 5261
QY 431 ACATCTAGAGCCTACATTAACCTGCTTTGTAGAGCCCGGAGAGAGATTTTCAACTGA 490
DB 5262 ACCTGGTGAACCTTCCCTTGACAGCTTTGACATGCCCGGAGAGAGATTTTCAACTGA 5321
QY 491 TGGAGAAAGGAT 501
DB 5322 TGGAGAAAGGAT 5332
XX
RESULT 9
ABL99976
ID ABL99976 standard; DNA; 378 BP.
XX
AC ABL99976;
XX
DT 14-AUG-2002 (first entry)
XX
DE Rat disease associated gene related polynucleotide SEQ ID NO 3.
XX
KW Rat; cardiac; heart disease; cardiovascular disease; cardiac infarction;
KW angina; gene therapy; ds.
XX
OS Rattus sp.
XX
FN WO200233082-A1.
XX
PD 25-APR-2002.
XX
PF 18-OCT-2001; 2001WO-JP09140.
XX
PR 19-OCT-2000; 2000JP-0319912.
XX
PR 16-NOV-2000; 2000JP-0350183.
XX
PA (TAKE ) TAKEDA CHEM IND LTD.
XX
PI Koyama N, Tanida S, Watanabe T;
XX
WPI; 2002-394557/42.
XX
Disease-associated gene and encoded RGS5-like protein, applicable in
diagnosis and prevention or treatment of heart diseases e.g.
cardiovascular diseases, cardiac infarction, heart failure and angina,
including gene therapy -
XX
Example 1; Page 73; 82pp; Japanese.
XX
The invention relates to a protein (ABB83788) or its salt. The protein
and encoded DNA (ABL99976) are applicable in diagnosis and prevention or
treatment of heart diseases e.g. cardiovascular diseases, cardiac
infarction, heart failure and angina, including gene therapy. The present
sequence is that of a polynucleotide, useful in examples of the
invention.
XX
Sequence 378 BP; 115 A; 101 C; 80 G; 82 T; 0 other;
SQ
```

```
Query Match
Best Local Similarity 9.8%; Score 49; DB 24; Length 378;
Matches 73; Conservative 0; Mismatches 40; Indels 0; Gaps 0;
XX
QY 389 AGGTGAACCTGGATTCTTGCACCGGGAGAGACAAGCCGGAACATGCTAGAGCTTACAA 448
DB 65 AGGTGAACATTTGACCACCTTCACTAAGACATCAACCATGAGAACCTGGTGAACCTTCCC 124
QY 449 TAACCTGTTGTAGAGGCCCGGAGAGAGATTTTCAACCTGATGGAGAGGAT 501
```



XX 14-AUG-2002 (first entry)  
DT Rat disease associated gene related polynucleotide SEQ ID NO 6.  
DE Rat; cardiact; heart disease; cardiovascular disease; cardiac infarction;  
KW angina; gene therapy; ds.  
XX  
OS Rattus sp.  
XX  
PN WO200233082-A1.  
XX  
PD 25-APR-2002.  
XX  
PF 18-OCT-2001; 2001WO-JP09140.  
XX  
PR 19-OCT-2000; 2000JP-0319912.  
XX  
PR 16-NOV-2000; 2000JP-0350183.  
XX  
PA (TAKE ) TAKEDA CHEM IND LTD.  
XX  
PI Koyama N, Tanida S, Watanabe T;  
XX  
DR WPI; 2002-394557/42.  
XX  
PT Disease-associated gene and encoded RGS5-like protein, applicable in  
PT diagnosis and prevention or treatment of heart diseases e.g.  
PT cardiovascular diseases, cardiac infarction, heart failure and angina,  
PT including gene therapy -  
XX  
PS Example 1; Page 74-75; 82pp; Japanese.  
XX  
CC The invention relates to a protein (ABB83788) or its salt. The protein  
CC and encoded DNA (ABU99975) are applicable in diagnosis and prevention or  
CC treatment of heart diseases e.g. cardiovascular diseases, cardiac  
CC infarction, heart failure and angina, including gene therapy. The present  
CC sequence is that of a polynucleotide, useful in examples of the  
CC invention.  
XX  
SQ Sequence 817 BP; 228 A; 214 C; 192 G; 183 T; 0 other;  
Query Match 9.8%; Score 49; DB 24; Length 817;  
Best Local Similarity 64.6%; Pred. No. 6.1e-05;  
Matches 73; Conservative 0; Mismatches 40; Indels 0; Gaps 0;  
QY 389 AGGTGAACCTGGATCTTGCACCGGAGAGACAGACAGCGGACATGCTAGAGCTACAA 448  
Db 504 AGGTGAACCTGGATCTTGCACCGGAGAGACAGACAGCGGACATGCTAGAGAGGAT 563  
QY 449 TAACCTGCTTTGATGAGGCCCGGAGAGAGATTTTCAACCTGATGAGAGGAT 501  
Db 564 CTCACAGCTTTGACCTGGCCCGGAGAGAGATTTACGCTGATGAGAGGAT 616  
RESULT 13  
ABA92505  
ID ABA92505 standard; cDNA; 540 BP.  
XX  
AC ABA92505;  
XX  
DT 19-MAR-2002 (first entry)  
XX  
DE Human regulator of G protein signalling (RGS8) encoding cDNA.  
DE  
KW Human; regulator of G protein signalling; RGS8; cerebroprotective;  
KW vulnerary; tranquilliser; analgesic; anticonvulsant; vasotropic;  
KW vaccine; stroke; head trauma; anxiety; pain; epileptic seizure; ss.  
XX  
OS Homo sapiens.  
XX  
FH Key Location/Qualifiers  
FT CDS 1..540  
/\*tag= a

FT  
FT /partial  
FT /product= "regulator of G protein signalling (RGS8)"  
XX /note= "no stop codon given"  
PN WO200185937-A2.  
XX  
XX 15-NOV-2001.  
XX  
PF 10-MAY-2001; 2001WO-EP05295.  
XX  
PR 12-MAY-2000; 2000EP-0110163.  
XX  
PA (MERE ) MERCK PATENT GMBH.  
XX  
PI Wilm C, Gassen M;  
XX  
DR WPI; 2002-055595/07.  
DR P-PSDB; ABB04999.  
XX  
PT Novel regulator of G-protein signaling polypeptide and polynucleotide  
PT for diagnosing, treating stroke, head trauma, anxiety, pain, epileptic  
PT seizures and for identifying modulators of therapeutic use -  
XX  
PS Claim 5; Page 34-35; 35pp; English.  
XX  
CC The present sequence encodes a human regulator of G-protein signalling,  
CC designated RGS8. RGS8 has cerebroprotective, vulnerary, tranquilliser,  
CC analgesic, anticonvulsant and vasotropic activities, and can be used in  
CC vaccine production. The RGS8 protein is useful in screening assays to  
CC identify compounds that stimulate or inhibit the function or level of  
CC the protein. RGS8 proteins and polynucleotides are useful as vaccines.  
CC The proteins are useful as immunogens to produce antibodies which are  
CC useful for treating diseases, to isolate or to identify clones  
CC expressing the protein or to purify the proteins by affinity  
CC chromatography. RGS8 proteins are also useful to identify membrane  
CC bound or soluble receptors. The RGS8 polynucleotide is useful for the  
CC recombinant production of RGS8 proteins, as hybridisation probes for  
CC cDNA and genomic DNA, as primers for nucleic acid amplification reaction  
CC to isolate full-length cDNAs and genomic clones encoding RGS8 proteins,  
CC in diagnostic assays by detecting mutations in the associated gene, for  
CC chromosome localisation studies, tissue expression studies and for  
CC producing transgenic animals useful in drug discovery and target  
CC validation. RGS8 proteins and polynucleotides can be used in the  
CC diagnosis and treatment of stroke, head trauma, anxiety, pain, epileptic  
CC seizures and for identifying modulators of therapeutic use.  
XX  
SQ Sequence 540 BP; 143 A; 131 C; 147 G; 119 T; 0 other;  
Query Match 9.6%; Score 48; DB 24; Length 540;  
Best Local Similarity 64.3%; Pred. No. 0.0001;  
Matches 72; Conservative 0; Mismatches 40; Indels 0; Gaps 0;  
QY 389 AGGTGAACCTGGATCTTGCACCGGAGAGACAGCGGACATGCTAGAGCTACAA 448  
Db 359 AGGTGAACCTGGATCTTGCACCGGAGAGACAGCGGACATGCTAGAGAGGAT 418  
QY 449 TAACCTGCTTTGATGAGGCCCGGAGAGAGATTTTCAACCTGATGAGAGGAT 500  
Db 419 TGACTTGTCTTGGACCAAGCCCGGAGAGAGATTTACAGCTCATGAGAGAGA 470  
RESULT 14  
AAA39678  
ID AAA39678 standard; cDNA; 606 BP.  
XX  
XX AAA39678;  
AC  
XX  
DT 18-SEP-2000 (first entry)  
XX  
DE Cytokine response gene-related cDNA sequence (Seq ID 27).  
XX  
KW CR2; human; antibody; cytokine response gene; cytostatic; anti-allergic;  
KW immunosuppressive; antimicrobial; therapy; cell proliferation; treatment;



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OM nucleic - nucleic search, using sw model

Run on: November 6, 2003, 21:13:51 ; Search time 155.728 Seconds  
(without alignments)  
8684.478 Million cell updates/sec

Title: US-09-939-209A-3\_COPY\_19800\_20300

Perfect score: 501

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Scoring table: IDENTITY\_NUC

Gapop 10.0 , Gapext 1.0

Searched: 2552756 seqs, 1349719017 residues

Total number of hits satisfying chosen parameters: 5105512

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

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23: /SIDSI/gcgdata/geneseq/geneseq-emb1/NA2001A.DAT.\*  
24: /SIDSI/gcgdata/geneseq/geneseq-emb1/NA2001B.DAT.\*  
25: /SIDSI/gcgdata/geneseq/geneseq-emb1/NA2002.DAT.\*  
26: /SIDSI/gcgdata/geneseq/geneseq-emb1/NA2003.DAT.\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

#### SUMMARIES

Result No.	Score	Query Match	Length	ID	Description
1	501	100.0	20300	24	ABK47337
2	479.4	95.7	4709	21	AAC69112
3	478.6	95.5	5065	21	AAC69111
4	478.6	95.5	5065	22	AAF33219
5	477	95.2	1494	23	AAS68523
6	477	95.2	3192	23	AAS79105
7	477	95.2	3192	23	ANS81387
8	477	95.2	3432	23	AAS68186

9	477	95.2	3432	23	AAS70808	DNA encoding novel
10	477	95.2	3432	23	AAS81679	DNA encoding novel
11	477	95.2	3434	23	AAS87390	DNA encoding novel
12	477	95.2	4137	23	AAS74444	DNA encoding novel
13	477	95.2	4137	23	AAS78954	DNA encoding novel
14	477	95.2	4137	23	AAS87865	DNA encoding novel
15	477	95.2	4202	23	AAS73338	DNA encoding novel
16	477	95.2	5367	23	AAS70936	DNA encoding novel
17	477	95.2	7130	23	AAS78741	DNA encoding novel
18	475.4	94.9	1785	23	AAS68655	DNA encoding novel
19	475.4	94.9	1785	23	AAS79132	DNA encoding novel
20	475.4	94.9	1919	23	AAS69603	DNA encoding novel
21	475.4	94.9	1944	23	AAS77252	DNA encoding novel
22	475.4	94.9	2025	23	AAS69612	DNA encoding novel
23	475.4	94.9	2025	23	AAS73299	DNA encoding novel
24	475.4	94.9	2025	23	AAS77455	DNA encoding novel
25	475.4	94.9	2025	23	AAS76700	DNA encoding novel
26	475.4	94.9	2025	23	AAS78928	DNA encoding novel
27	475.4	94.9	2107	23	AAS74303	DNA encoding novel
28	475.4	94.9	3558	23	AAS72863	DNA encoding novel
29	475.4	94.9	3562	23	AAS79127	DNA encoding novel
30	475.4	94.9	3657	23	AAS70241	DNA encoding novel
31	475.4	94.9	3657	23	AAS73824	DNA encoding novel
32	475.4	94.9	3657	23	AAS74283	DNA encoding novel
33	475.4	94.9	3657	23	AAS74983	DNA encoding novel
34	475.4	94.9	3657	23	AAS78725	DNA encoding novel
35	475.4	94.9	3657	23	AAS78940	DNA encoding novel
36	475.4	94.9	3657	23	AAS81661	DNA encoding novel
37	475.4	94.9	3657	23	AAS84092	DNA encoding novel
38	475.4	94.9	4251	23	AAS68182	DNA encoding novel
39	475.4	94.9	4555	23	AAS84104	DNA encoding novel
40	475.4	94.9	4797	23	AAS81681	DNA encoding novel
41	475.4	94.9	5215	23	AAS70816	DNA encoding novel
42	475.4	94.9	6194	23	AAS73339	DNA encoding novel
43	475.4	94.9	13234	23	AAS82685	DNA encoding novel
44	475.4	94.9	1503841	24	ABT00010	Human neuregulin 1
45	475.4	94.9	1503841	24	ABT01503	Human neuregulin 1

#### ALIGNMENTS

##### RESULT 1

ABK47337  
ID ABK47337 standard; DNA; 20300 BP.

XX AC ABK47337;

XX DT 18-JUN-2002 (first entry)

XX DE Genomic nucleotide sequence encoding human RGS-4 protein.

XX DE RGS-4; schizophrenia; human; regulator of G protein signalling 4;  
XX KW neuroleptic; polymorphism; schizotypy; schizoid; chromosome 1q21-22;  
XX KW gene; ds.

XX OS Homo sapiens.

XX FH Key

FT variation Location/Qualifiers

FT replace (4121,T)

FT /\*tag= a

FT FT standard name= "Single-nucleotide polymorphism"

FT replace (4123,A)

FT /\*tag= b

FT FT standard name= "Single-nucleotide polymorphism"

FT replace (4368,C)

FT /\*tag= c

FT FT standard name= "Single-nucleotide polymorphism"

FT replace (4621,C)

FT /\*tag= d

FT FT standard name= "Single-nucleotide polymorphism"

FT replace (4790,T)

FT /\*tag= e

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FT replace (4816,T)  
FT /tag= f  
FT /standard_name= "Single-nucleotide polymorphism"  
FT replace (4970,T)  
FT /tag= g  
FT /standard_name= "Single-nucleotide polymorphism"  
FT replace (5055,G)  
FT /tag= h  
FT /standard_name= "Single-nucleotide polymorphism"  
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FT replace (5695,A)  
FT /tag= j  
FT /standard_name= "Single-nucleotide polymorphism"  
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FT /tag= l  
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FT /note= "Deletion polymorphism"  
FT replace (9892,A)  
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FT /standard_name= "Single-nucleotide polymorphism"  
FT replace (12145,T)  
FT /tag= w  
FT /standard_name= "Single-nucleotide polymorphism"  
FT replace (14367,G)  
FT /tag= x  
FT /standard_name= "Single-nucleotide polymorphism"  
FT replace (17027..17029,GA)  
FT /tag= y  
FT /note= "Deletion polymorphism"  
FT replace (17630,T)  
FT /tag= z  
FT /standard_name= "Single-nucleotide polymorphism"  
XX  
PN WO200216653-A2.  
XX  
PD 28-FEB-2002.  
XX  
PF 24-AUG-2001; 2001WO-US26622.  
XX  
PR 24-AUG-2000; 2000US-228021P.  
XX
```

(UYPI-) UNIV PITTSBURGH.  
Levitt PR, Mirnics K, Kodavali VC, Nimgaonkar VL;  
WPI; 2002-292070/33.  
Diagnosing, assessing susceptibility and treating schizophrenia,  
PT involves observing regulator of G-protein signalling 4, RGS4 levels in a  
XX subject -  
PS Claim 1; Page 20-33; 112pp; English.  
XX  
CC This invention relates to a novel method for diagnosing schizophrenia  
or determining susceptibility to schizophrenia in a human. The method  
comprises obtaining from a patient a DNA sample and detecting variations  
in the regulator of G-protein signalling 4 (RGS4) gene. Alternatively,  
CC the method involves measuring RGS4 mRNA or protein levels in a tissue  
sample from the patient and determining if there is a reduced level.  
CC The method of the invention is useful for diagnosing and determining  
susceptibility to schizophrenia. The invention also comprises a method  
that is useful for treating schizophrenia which includes a prophylactic  
CC treatment. The method of genotyping polymorphic variants in the RGS-4  
gene is applied to diagnosing pathologies of the schizophrenic spectrum,  
CC such as in particular schizotypy, schizoid individuals, etc. This  
method offers the possibility of diagnosing schizophrenia by a  
CC biological test and no longer exclusively by clinical evaluations.  
CC The present sequence represents the genomic DNA encoding the human  
regulator of G-protein signalling 4 (RGS4) protein used in the method of  
CC the invention. The gene for the RGS4 protein is located on human  
CC chromosome 1q21-22.  
XX  
SQ Sequence 20300 BP; 6157 A; 4102 C; 3775 G; 6266 T; 0 other;  
Query Match 100.0%; Score 501; DB 24; Length 20300;  
Best Local Similarity 100.0%; Pred. No. 3e-133;  
Matches 501; Conservative 0; Mismatches 0; Indels 0; Gaps 0;  
QY 1 CCACATGATTATCTCAATAGATGCGAGAAAGGCAATTGACAAAATTTAAACAATCTTTTCAT 60  
DB 19800 CCACATGATTATCTCAATAGATGCGAGAAAGGCAATTGACAAAATTTAAACAATCTTTTCAT 19859  
QY 61 GCTAAACTCTCAATCAATAGTATGATGGAGCGTATCTCAAAATTAAGCACTAT 120  
DB 19860 GCTAAACTCTCAATCAATAGTATGATGGAGCGTATCTCAAAATTAAGCACTAT 19919  
QY 121 CTATGACAACTCAGAGCAATATCATCTGAAATGGGCAAAACTGGAAGCATTCCTTT 180  
DB 19920 CTATGACAACTCAGAGCAATATCATCTGAAATGGGCAAAACTGGAAGCATTCCTTT 19979  
QY 181 GAAACGGGCAACAGACAGGGATGCCCTCTCTCACACTCTCTTCAACATAGTGTGGA 240  
DB 19980 GAAACGGGCAACAGACAGGGATGCCCTCTCTCACACTCTCTTCAACATAGTGTGGA 20039  
QY 241 AGCTCTGCCAGGCAATTAGGAGGAGGAAGGAATTAAGGCTATCAATTAGGAGAGA 300  
DB 20040 AGCTCTGCCAGGCAATTAGGAGGAGGAAGGAATTAAGGCTATCAATTAGGAGAGA 20099  
QY 301 GGAAGTCAAAATGTCTCTGTTGTCAGATGACATGATTTGATATCTAGAAAACCCATCGT 360  
DB 20100 GGAAGTCAAAATGTCTCTGTTGTCAGATGACATGATTTGATATCTAGAAAACCCATCGT 20159  
QY 361 CTCAGCCCAAAATCTCTTAAGCTGATAAGCAACTTCAGCAAGTCTCAGGATACAAAT 420  
DB 20160 CTCAGCCCAAAATCTCTTAAGCTGATAAGCAACTTCAGCAAGTCTCAGGATACAAAT 20219  
QY 421 CAATGTACAAAATTCACAGCACTCTTATATCATCAATTAACAGCAAAACAGAGAGCAAT 480  
DB 20220 CAATGTACAAAATTCACAGCACTCTTATATCATCAATTAACAGCAAAACAGAGAGCAAT 20279  
QY 481 CATGAGTCAACTCCCATTCAC 501  
DB 20280 CATGAGTCAACTCCCATTCAC 20300





CC generate fusion proteins by linking to the gene for the human  
 CC immunoglobulin G Fc portion (AAC69075) for increasing the stability of  
 CC the fusion protein as compared to the human protein only. The genes and  
 CC proteins are useful for preventing, ameliorating or treating medical  
 CC conditions, e.g. by protein or gene therapy. The genes are isolated  
 CC from a range of human tissues disclosed in the specification. The  
 CC nucleic acids, proteins, antibodies and (ant)agonists are useful in  
 CC the diagnosis, treatment and prevention of: (a) cancer, e.g. breast  
 CC and ovarian cancer, and other cancers of the adrenal gland, bone, bone  
 CC marrow, breast, gastrointestinal tract, liver, lung, or urogenital;  
 CC (b) immune disorders e.g. Addison's disease, allergies, autoimmune  
 CC haemolytic anaemia, autoimmune thyroiditis, diabetes mellitus, Crohn's  
 CC disease, multiple sclerosis, rheumatoid arthritis and ulcerative  
 CC colitis; (c) cardiovascular disorders such as myocardial ischaemia; (d)  
 CC wound healing; (e) neurological diseases e.g. cerebral anoxia and  
 CC epilepsy; and (f) infectious diseases such as viral, bacterial, fungal  
 CC and parasitic infections.

XX Sequence 5065 BP; 980 A; 948 C; 1046 G; 2017 T; 74 other;

Query Match 95.5%; Score 478.6; DB 21; Length 5065;  
 Best Local Similarity 96.6%; Pred. No. 4.8e-127;  
 Matches 484; Conservative 4; Mismatches 13; Indels 0; Gaps 0;

QY 1 CCACATGATTATCTCAATAGATGCGAAGAGGCATTGACAAAATTACAACTTTCAT 60  
 DB 2312 CCACATGATTATCTCAATAGATGCGAAGAGGCATTGACAAAATTACAACTTTCAT 2253  
 QY 61 GCTAAAACTCTCAATCAATAGGATTTGATGGAGCTATCTCAAAATAATAGCACTAT 120  
 DB 2252 GCTAAAACTCTCAATCAATAGGATTTGATGGAGCTATCTCAAAATAATAGCACTAT 2193  
 QY 121 CTATGACAAACTCAGCGCAATATCATCTGAAATGGCGAAAACTGGAAAGCAATCCCTTT 180  
 DB 2192 CTATGACAAACTCAGCGCAATATCATCTGAAATGGCGAAAACTGGAAAGCAATCCCTTT 2133  
 QY 181 GAAAAAGGGGCAAGACAGGGATGCCCTCTCCACACCTCCTATTCAACATAGTGTGA 240  
 DB 2132 GAAAAAGGGGCAAGACAGGGATGCCCTCTCCACACCTCCTATTCAACATAGTGTGA 2073  
 QY 241 AGCTCTGGCCAGGGCAATPAGGACGAGAGGAAATAAGGGTATTCAATTAGGAGAGA 300  
 DB 2072 AGTTCTGGCCAGGGCAATPAGGACGAGAGGAAATAAGGGTATTCAATTAGGAGAGA 2013  
 QY 301 GGAAGTCAAAATGTCCTGTTTGCAGATCATGATGTTATATCTAGAAAAACCCCATCGT 360  
 DB 2012 GGAAGTCAAAATGTCCTGTTTGCAGATCATGATGTTATATCTAGAAAAACCCCATG 1953  
 QY 361 CTCAGCCCAAAATCTCCTTAAGCTGATAGCAACTTCAGCAAGTCTCAGGATACAAAT 420  
 DB 1952 CTCAGCCCAAAATCTCCTTAAGCTGATAGCAACTTCAGCAAGTCTCAGGATACAAAT 1893  
 QY 421 CAATGTCAAAAATCAGAAAGCTCTTATACATCAATAACAGACAAACAGAGAGCAAAAT 480  
 DB 1892 CAATGTCAAAAATCAGAAAGCTCTTATACCAAAVAACAGACAAACAGAGAGCAAAAT 1833  
 QY 481 CATGAGTGAATCCCATTCAC 501  
 DB 1832 CATGAGTGAATCCCATTCAC 1812

## RESULT 4

AAF33219/c

ID AAF33219 standard; cDNA; 5065 BP.

XX AAF33219;

AC AAF33219;

XX 23-MAR-2001 (first entry)

XX Human secreted protein gene 7 SEQ ID NO:17.

XX Human; secreted protein; diagnosis; immunomodulatory; antisclerotic;

XX dermatological; immunosuppressive; antiinflammatory; anti-HIV;

KW

immunostimulant; cytostatic; cardiant; vascular; anti-angiogenic;

ophthalmological; neuroprotectant; nootropic; anticonvulsant; vulnary;

antialzheimers; antiparkinsonian; antimicrobial; immune disorder;

multiple sclerosis; systemic lupus erythematosus; HIV; infection;

hyperproliferative disorder; cancer; Gaucher's disease; wound healing;

cardiovascular disease; Scimitar syndrome; Chaga's cardiomyopathy;

coronary arteriosclerosis; angiogenic disorder; diabetic retinopathy;

corneal graft neovascularisation; neurodegenerative disorder; regeneration;

Huntington's chorea; Alzheimer's disease; Parkinson's disease;

infectious disease; chemotaxis; ss.

XX Homo sapiens.

OS

XX WO2000076530-A1.

XX 21-DEC-2000.

XX 01-JUN-2000; 2000WO-US14933.

XX 11-JUN-1999; 99US-0138572.

XX (HUMA-) HUMAN GENOME SCI INC.

XX (ROSE/) ROSEN C A.

XX Rosen CA, Ruben SM, Komatsoulis GA;

XX WPI; 2001-071147/08.

XX P-PSDB; AAB64888.

XX Nucleic acids encoding 49 human secreted polypeptides, useful for

preventing, diagnosing and/or treating e.g. cancers, Parkinson's

disease and diabetic retinopathy -

Claim 1; Page 453-455; 554pp; English.

XX The polynucleotide sequences given in AAF33213 to AAF33261 encode the

human secreted proteins given in AAB64882 to AAB64930. AAB64931 to

AAB64991 represent human secreted polypeptide sequences and proteins

homologous to them, which are given in the exemplification of the present

invention. Human secreted proteins have activities based on the tissues

and cells the genes are expressed in. Examples of activities include:

immunomodulatory; antisclerotic; dermatological; immunosuppressive;

antiflammatory; anti-HIV; immunostimulant; cytostatic; Cardiant;

vascular; antimicrobial; anti-angiogenic; ophthalmological;

antiparkinsonian; anticonvulsant; nootropic; antialzheimers;

be used in the prevention, diagnosis and treatment of diseases associated

with inappropriate polypeptide expression. Disorders that may be

prevented, diagnosed and/or treated by the above methods include immune

disorders (e.g. multiple sclerosis, systemic lupus erythematosus and

human immuno-deficiency virus (HIV) infections), hyperproliferative

disorders (e.g. cancers and Gaucher's disease), cardiovascular diseases

(e.g. Scimitar syndrome, Chaga's cardiomyopathy and coronary

arteriosclerosis), angiogenic disorders (e.g. corneal graft

neovascularisation and diabetic retinopathy), neurological disorders

(e.g. Huntington's chorea, Alzheimer's disease and Parkinson's disease),

infectious diseases and/or for promoting wound healing, regeneration and

/or chemotaxis. AAF33204 to AAF33212 and AAB64881 represent sequences

used in the exemplification of the present invention.

XX Sequence 5065 BP; 980 A; 948 C; 1046 G; 2017 T; 74 other;

Query Match 95.5%; Score 478.6; DB 22; Length 5065;

Best Local Similarity 96.6%; Pred. No. 4.8e-127;

Matches 484; Conservative 4; Mismatches 13; Indels 0; Gaps 0;

QY 1 CCACATGATTATCTCAATAGATGCGAAGAGGCATTGACAAAATTACAACTTTCAT 60

DB 2312 CCACATGATTATCTCAATAGATGCGAAGAGGCATTGACAAAATTACAACTTTCAT 2253

QY 61 GCTAAAACTCTCAATCAATAGGATTTGATGGAGCTATCTCAAAATAATAGCACTAT 120

DB 2252 GCTAAAACTCTCAATCAATAGGATTTGATGGAGCTATCTCAAAATAATAGCACTAT 2193



OS Homo sapiens.  
XX WO200175067-A2.  
XX PD 11-OCT-2001.  
XX PF 30-MAR-2001; 2001WO-US08631.  
XX PR 31-MAR-2000; 2000US-0540217.  
XX PR 23-AUG-2000; 2000US-0649167.  
XX PA (HYSE-) HYSEQ INC.  
XX PI Drmanac RT, Liu C, Tang YT;  
XX DR WPI; 2001-639362/73.  
XX DR P-PSDB; ABG14918.  
XX PT New isolated polynucleotide and encoded polypeptides, useful in  
XX PT diagnostics, forensics, gene mapping, identification of mutations  
XX PT responsible for genetic disorders or other traits and to assess  
XX PT biodiversity -  
XX PS Claim 1; SEQ ID No 14909; 103pp; English.  
XX CC The invention relates to isolated polynucleotide (I) and  
XX CC polypeptide (II) sequences. (I) is useful as hybridisation probes,  
XX CC polymerase chain reaction (PCR) primers, oligomers, and for chromosome  
XX CC and gene mapping, and in recombinant production of (II). The  
XX CC polynucleotides are also used in diagnostics as expressed sequence tags  
XX CC for identifying expressed genes. (I) is useful in gene therapy techniques  
XX CC to restore normal activity of (II) or to treat disease states involving  
XX CC (II). (II) is useful for generating antibodies against it, detecting or  
XX CC quantitating a polypeptide in tissue, as molecular weight markers and as  
XX CC a food supplement. (II) and its binding partners are useful in medical  
XX CC disorders involving aberrant protein expression or biological activity.  
XX CC The polypeptide and polynucleotide sequences have applications in  
XX CC diagnostics, forensics, gene mapping, identification of mutations  
XX CC responsible for genetic disorders or other traits to assess biodiversity  
XX CC and to produce other types of data and products dependent on DNA and  
XX CC amino acid sequences. AAS64197-AAS94564 represent novel human  
XX CC diagnostic coding sequences of the invention.  
XX CC Note: The sequence data for this patent did not appear in the printed  
XX CC specification, but was obtained in electronic format directly from WIPO  
XX CC at ftp.wipo.int/pub/published\_pct\_sequences.  
XX SQ Sequence 3192 BP; 1290 A; 705 C; 562 G; 635 T; 0 other;  
  
Query Match 95.2%; Score 477; DB 23; Length 3192;  
Best Local Similarity 97.0%; Pred. No. 1.2e-126;  
Matches 486; Conservative 0; Mismatches 15; Indels 0; Gaps 0;  
  
QY 1 CCACATGATTATCTCAATAGATGCGAGAAAGGCATTTCACAAATTTAACCACTCTTCAT 60  
Db 1353 CCACATGATTATCTCAATAGATGCGAGAAAGGCATTTCACAAATTTACCACTCTTCAT 1412  
QY 61 GCTAAAACTCTCAATCAATAGGATTTGATGGGACGTATCTCAAAATAAAGCACTAT 120  
Db 1413 GCTAAAACTCTCAATCAATAGGATTTGATGGGACGTATCTCAAAATAAAGCACTAT 1472  
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Db 1473 CTATGACAACTCACAGCAATATCATCTGATGGGCAAAACCTGGAGCAATTCCTTT 1532  
QY 181 GAAACGGGCAACAGACGGGATGCCCTCTCTCACCTCTCTTATTCACATAGTTTGA 240  
Db 1533 GAAACCTGGCAACAGACGGGATGCCCTCTCTCACCTCTCTTATTCACATAGTTTGA 1592  
QY 241 AGCTTGGCCAGGCAATTAGCAGAGAGAGAAATAAGGTTATCAATAGGAGAGA 300  
Db 1593 AGTCTGGCCAGGCAATTAGCAGAGAGAGAAATAAGGTTATCAATAGGAGAGA 1652

QY 301 GGAAGTCAAATTTGCTCCCTGTTTGCAGATGACATGATTGTATATCTAGAAAAACCCCATCGT 360  
Db 1653 GGAAGTCAAATTTGCTCCCTGTTTGCAGATGACATGATTGTATATCTAGAAAAACCCCATTTGT 1712  
QY 361 CTCAGCCCAAAATCTCTTAAGCTGATTAAGCAACTTCAGCAAGTCTCAGGATACAAAAT 420  
Db 1713 CTCAGCCCAAAATCTCTTAAGCTGATTAAGCAACTTCAGCAAGTCTCAGGATACAAAAT 1772  
QY 421 CAATGTACAAAAATCACAGCACTCTTATACATCAATAACAGCAAAACAGAGAGCCAAAT 480  
Db 1773 CAATGTACAAAAATCACAGCACTCTTATACATCAATAACAGCAAAACAGAGAGCCAAAT 1832  
QY 481 CATGAGTGAACCTCCCATTCAC 501  
Db 1833 CATGAGTGAACCTCCCATTCAC 1853  
  
RESULT 7  
AAS81387  
ID AAS81387 standard; cDNA; 3192 BP.  
XX AC AAS81387;  
XX DT 13-FEB-2002 (first entry)  
XX DE DNA encoding novel human diagnostic protein #17191.  
XX KW Human; chromosome mapping; gene mapping; gene therapy; forensic;  
XX KW food supplement; medical imaging; diagnostic; genetic disorder; ss.  
XX OS Homo sapiens.  
XX PN WO200175067-A2.  
XX PD 11-OCT-2001.  
XX PF 30-MAR-2001; 2001WO-US08631.  
XX PR 31-MAR-2000; 2000US-0540217.  
XX PR 23-AUG-2000; 2000US-0649167.  
XX PA (HYSE-) HYSEQ INC.  
XX PI Drmanac RT, Liu C, Tang YT;  
XX DR WPI; 2001-639362/73.  
XX DR P-PSDB; ABG17200.  
XX PT New isolated polynucleotide and encoded polypeptides, useful in  
XX PT diagnostics, forensics, gene mapping, identification of mutations  
XX PT responsible for genetic disorders or other traits and to assess  
XX PT biodiversity -  
XX PS Claim 1; SEQ ID No 17191; 103pp; English.  
XX CC The invention relates to isolated polynucleotide (I) and  
XX CC polypeptide (II) sequences. (I) is useful as hybridisation probes,  
XX CC polymerase chain reaction (PCR) primers, oligomers, and for chromosome  
XX CC and gene mapping, and in recombinant production of (II). The  
XX CC polynucleotides are also used in diagnostics as expressed sequence tags  
XX CC for identifying expressed genes. (I) is useful in gene therapy techniques  
XX CC to restore normal activity of (II) or to treat disease states involving  
XX CC (II). (II) is useful for generating antibodies against it, detecting or  
XX CC quantitating a polypeptide in tissue, as molecular weight markers and as  
XX CC a food supplement. (II) and its binding partners are useful in medical  
XX CC disorders involving aberrant protein expression or biological activity.  
XX CC The polypeptide and polynucleotide sequences have applications in  
XX CC diagnostics, forensics, gene mapping, identification of mutations  
XX CC responsible for genetic disorders or other traits to assess biodiversity  
XX CC and to produce other types of data and products dependent on DNA and  
XX CC amino acid sequences. AAS64197-AAS94564 represent novel human  
XX CC diagnostic coding sequences of the invention.  
XX CC Note: The sequence data for this patent did not appear in the printed  
XX CC specification, but was obtained in electronic format directly from WIPO  
XX CC at ftp.wipo.int/pub/published\_pct\_sequences.  
XX SQ Sequence 3192 BP; 1290 A; 705 C; 562 G; 635 T; 0 other;



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Db      1833 CATGAGTGAACCTCCCATTCAC 1853
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RESULT 9
AAS70808
ID AAS70808 standard; cDNA; 3432 BP.
XX
AC AAS70808;
XX
DT 13-FEB-2002 (first entry)
XX
DE DNA encoding novel human diagnostic protein #6612.
XX
KW Human; chromosome mapping; gene mapping; gene therapy; forensic;
KW food supplement; medical imaging; diagnostic; genetic disorder; ss.
XX
OS Homo sapiens.
XX
PN WO200175067-A2.
XX
PD 11-OCT-2001.
XX
PF 30-MAR-2001; 2001WO-US08631.
XX
PR 31-MAR-2000; 2000US-0540217.
PR 23-AUG-2000; 2000US-0649167.
XX
PA (HYSE-) HYSEQ INC.
XX
PI Drmanac RT, Liu C, Tang YT;
XX
DR WPI; 2001-639362/73.
DR P-PSDB; ABG06621.
XX
XX
New isolated polynucleotide and encoded polypeptides, useful in
PT diagnostics, forensics, gene mapping, identification of mutations
PT responsible for genetic disorders or other traits and to assess
PT biodiversity -
XX
PS Claim 1; SEQ ID No 6612; 103pp; English.
XX
XX
The invention relates to isolated polynucleotide (I) and
XX polypeptide (II) sequences. (I) is useful as hybridization probes,
XX polymerase chain reaction (PCR) primers, oligomers, and for chromosome
XX and gene mapping, and in recombinant production of (II). The
XX polynucleotides are also used in diagnostics as expressed sequence tags
XX for identifying expressed genes. (I) is useful in gene therapy techniques
XX to restore normal activity of (II) or to treat disease states involving
XX (II). (II) is useful for generating antibodies against it, detecting or
XX quantitating a polypeptide in tissue, as molecular weight markers and as
XX a food supplement. (II) and its binding partners are useful in medical
XX imaging of sites expressing (II). (I) and (II) are useful for treating
XX disorders involving aberrant protein expression or biological activity.
XX The polypeptide and polynucleotide sequences have applications in
XX diagnostics, forensics, gene mapping, identification of mutations
XX responsible for genetic disorders or other traits to assess biodiversity
XX and to produce other types of data and products dependent on DNA and
XX amino acid sequences. AAS64197-AAS94564 represent novel human
XX diagnostic coding sequences of the invention.
XX Note: The sequence data for this patent did not appear in the printed
XX specification, but was obtained in electronic format directly from WIPO
XX at ftp.wipo.int/pub/published_pct_sequences.
XX
SQ Sequence 3432 BP; 1349 A; 769 C; 624 G; 690 T; 0 other;

Query Match          95.2%; Score 477; DB 23; Length 3432;
Best Local Similarity 97.0%; Pred. No. 1.2e-126;
Matches 486; Conservative 0; Mismatches 15; Indels 0; Gaps 0;

QY      1 CCACATGATTATCTCAATAGTGCAGAAAAGCGATTTCGACAAAATTTAACAACTCTTCAT 60
|||||
Db      1353 CCACATGATTATCTCAATAGTGCAGAAAAGCGATTTCGACAAAATTTAACAACTCTTCAT 1412
|||||
QY      61 GCTAAAAAAGCTCTCAATCAATAGTGCAGCGTATTCATGCGAGCTATCTCAAAATAATAAGCACTAT 120
|||||
Db      1413 GCTAAAAAAGCTCTCAATCAATAGTGCAGCGTATTCATGCGAGCTATCTCAAAATAATAAGCACTAT 1472
|||||
QY      121 CTATGACAAAAGCTCAGAGCCAAATATCATACTGAATGGGCAAAAAGCTGGAAGCAATTCCTCTTT 180
|||||
Db      1473 CTATGACAAAAGCTCAGAGCCAAATATCATACTGAATGGGCAAAAAGCTGGAAGCAATTCCTCTTT 1532
|||||
QY      181 GAAAAAGGGGCAAGACAGAGGGGATGCCCTCTCTCACCACCTCCCTATTCAACATAGTGTGGA 240
|||||
Db      1533 GAAAAAGGGGCAAGACAGAGGGGATGCCCTCTCTCACCACCTCCCTATTCAACATAGTGTGGA 1592
|||||
QY      241 AGCTCTGGCCAGGGCAATTAGGCAAGAGGAATAAAGGGTATTCAATTAGGAGGAAGA 300
|||||
Db      1593 AGTTCTGGCCAGGGCAATTAGGCAAGAGGAATAAAGGGTATTCAATTAGGAGGAAGA 1652
|||||
QY      301 GGAAGTCAAAATTTGCTCCCTGTTTGCAGATGACATGATTTGATATCTAGAAAACCCCATCGT 360
|||||
Db      1653 GGAAGTCAAAATTTGCTCCCTGTTTGCAGATGACATGATTTGATATCTAGAAAACCCCATCGT 1712
|||||
QY      361 CTCAGCCCAAAATCTCCTTAAGCTGATAAGCAACTTCAGCAAAAGTCTCAGGATACAAAT 420
|||||
Db      1713 CTCAGCCCAAAATCTCCTTAAGCTGATAAGCAACTTCAGCAAAAGTCTCAGGATACAAAT 1772
|||||
QY      421 CAATGTACAAAATCACAAGCACTCTTATATACATCAATTAACAGACAAAACAGAGAGCCAAAT 480
|||||
Db      1773 CAATGTACAAAATCACAAGCACTCTTATATACCAACCAACAGACAAAACAGAGAGCCAAAT 1832
|||||
QY      481 CATGAGTGAACCTCCCATTCAC 501
Db      1833 CATGAGTGAACCTCCCATTCAC 1853
|||||

RESULT 10
AAS81679
ID AAS81679 standard; cDNA; 3432 BP.
XX
AC AAS81679;
XX
DT 13-FEB-2002 (first entry)
XX
DE DNA encoding novel human diagnostic protein #17483.
XX
KW Human; chromosome mapping; gene mapping; gene therapy; forensic;
KW food supplement; medical imaging; diagnostic; genetic disorder; ss.
XX
OS Homo sapiens.
XX
PN WO200175067-A2.
XX
PD 11-OCT-2001.
XX
PF 30-MAR-2001; 2001WO-US08631.
XX
PR 31-MAR-2000; 2000US-0540217.
PR 23-AUG-2000; 2000US-0649167.
XX
PA (HYSE-) HYSEQ INC.
XX
PI Drmanac RT, Liu C, Tang YT;
XX
DR WPI; 2001-639362/73.
DR P-PSDB; ABG17492.
XX
XX
New isolated polynucleotide and encoded polypeptides, useful in
PT diagnostics, forensics, gene mapping, identification of mutations
PT responsible for genetic disorders or other traits and to assess
PT biodiversity -
XX
PS Claim 1; SEQ ID No 17483; 103pp; English.
XX
XX
The invention relates to isolated polynucleotide (I) and

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QY 1 CCACATGATTATCTCAATAGATGCGAGAAAGGCATTTGACAAAATTTTAACTCTTTCAT 60  
Db |||||  
2265 CCACATGATTATCTCAATAGATGCGAGAAAGGCCTTTGACAAAATTTTAACTCTTTCAT 2324  
QY 61 GCTAAATCTCAATCAATTAAGTATTTGATGGAGCTATCTCAAAATTAAGCACTAT 120  
Db |||||  
2325 GCTAAATCTCAATCAATTAAGTATTTGATGGAGCTATCTCAAAATTAAGCACTAT 2384  
QY 121 CTATGACAACTCACAGGCAATATCATCTGAATGGGCAAAATCTGGAGCAATTCCTTT 180  
Db |||||  
2385 CTATGACAACTCACAGGCAATATCATCTGAATGGGCAAAATCTGGAGCAATTCCTTT 2444  
QY 181 GAAACGGGCAAGACAGGATGCTCTCTCACACATCTCTTATCAACATAGTGTGA 240  
Db |||||  
2445 GAAACGGGCAAGACAGGATGCTCTCTCACACATCTCTTATCAACATAGTGTGA 2504  
QY 241 AGCTCTGGCCAGGCAATPAGGACGAGAGGAATTAAGGCTATCAATTAAGGAGAGA 300  
Db |||||  
2505 AGTCTGGCCAGGCAATPAGGACGAGAGGAATTAAGGCTATCAATTAAGGAGAGA 2564  
QY 301 GGAAGTCAAAATGTCCTTTGCGATGACATGATTTGATATCTAGAAAACCCCATCT 360  
Db |||||  
2565 GGAAGTCAAAATGTCCTTTGCGATGACATGATTTGATATCTAGAAAACCCCATCT 2624  
QY 361 CTCAGGCCAAATCTCTTATGCTGATAGCAACTTCAGCAAGCTCTAGGATACAAAT 420  
Db |||||  
2625 CTCAGGCCAAATCTCTTATGCTGATAGCAACTTCAGCAAGCTCTAGGATACAAAT 2684  
QY 421 CAATGTACAAAATCACAGCACTCTTATACATCAATTAAGGCTATCAATTAAGGAGAGA 480  
Db |||||  
2685 CAATGTACAAAATCACAGCACTCTTATACATCAATTAAGGCTATCAATTAAGGAGAGA 2744  
QY 481 CATGAGTGAATCTCCCATTCAC 501  
Db |||||  
2745 CATGAGTGAATCTCCCATTCAC 2765

## RESULT 15

AA573338  
ID AAS73338 standard; cdna; 4202 BP.

XX AC AAS73338;  
XX

DT 13-FEB-2002 (first entry)

DE DNA encoding novel human diagnostic protein #9142.

XX Human; chromosome mapping; gene mapping; gene therapy; forensic;  
KW food supplement; medical imaging; diagnostic; genetic disorder; ss.

XX Homo sapiens.

XX WO200175067-A2.

XX 11-OCT-2001.

XX 30-MAR-2001; 2001WO-US08631.

XX 31-MAR-2000; 2000US-0540217.

XX 23-AUG-2000; 2000US-0649167.

XX (HYSE-) HYSEQ INC.

XX Drmanac RT, Liu C, Tang YT;

XX WPI; 2001-639362/73.

XX P-PSDB; ABG09151.

PT New isolated polynucleotide and encoded polypeptides, useful in  
PT diagnostics, forensics, gene mapping, identification of mutations  
PT responsible for genetic disorders or other traits and to assess  
PT biodiversity

XX

PS Claim 1; SEQ ID No 9142; 103pp; English.

CC The invention relates to isolated polynucleotide (I) and  
CC polypeptide (II) sequences. (I) is useful as hybridisation probes,  
CC polymerase chain reaction (PCR) primers, oligomers, and for chromosome  
CC and gene mapping, and in recombinant production of (II). The  
CC polynucleotides are also used in diagnostics as expressed sequence tags  
CC for identifying expressed genes. (I) is useful in gene therapy techniques  
CC to restore normal activity of (II) or to treat disease states involving  
CC (II). (II) is useful for generating antibodies against it, detecting or  
CC quantitating a polypeptide in tissue, as molecular weight markers and as  
CC a food supplement. (II) and its binding partners are useful in medical  
CC imaging of sites expressing (II). (I) and (II) are useful for treating  
CC disorders involving aberrant protein expression or biological activity.  
CC The polypeptide and polynucleotide sequences have applications in  
CC diagnostics, forensics, gene mapping, identification of mutations  
CC responsible for genetic disorders or other traits to assess biodiversity  
CC and to produce other types of data and products dependent on DNA and  
CC amino acid sequences. AAS64197-AAS94564 represent novel human  
CC diagnostic coding sequences of the invention.  
CC Note: The sequence data for this patent did not appear in the printed  
CC specification, but was obtained in electronic format directly from WIPO  
CC at ftp.wipo.int/pub/published\_pct\_sequences.

XX Sequence 4202 BP; 1482 A; 951 C; 928 G; 841 T; 0 other;

Query Match 95.2%; Score 477; DB 23; Length 4202;  
Best Local Similarity 97.0%; Pred. No. 1.3e-126;  
Matches 486; Conservative 0; Mismatches 15; Indels 0; Gaps 0;

QY 1 CCACATGATTATCTCAATAGATGCGAGAAAGGCATTTGACAAAATTTTAACTCTTTCAT 60

Db 2265 CCACATGATTATCTCAATAGATGCGAGAAAGGCCTTTGACAAAATTTTAACTCTTTCAT 2324

QY 61 GCTAAATCTCAATCAATTAAGTATTTGATGGAGCTATCTCAAAATTAAGCACTAT 120

Db 2325 GCTAAATCTCAATCAATTAAGTATTTGATGGAGCTATCTCAAAATTAAGCACTAT 2384

QY 121 CTATGACAACTCACAGGCAATATCATCTGAATGGGCAAAATCTGGAGCAATTCCTTT 180

Db 2385 CTATGACAACTCACAGGCAATATCATCTGAATGGGCAAAATCTGGAGCAATTCCTTT 2444

QY 181 GAAACGGGCAAGACAGGATGCTCTCTCACACATCTCTTATCAACATAGTGTGA 240

Db 2445 GAAACGGGCAAGACAGGATGCTCTCTCACACATCTCTTATCAACATAGTGTGA 2504

QY 241 AGCTCTGGCCAGGCAATPAGGACGAGAGGAATTAAGGCTATCAATTAAGGAGAGA 300

Db 2505 AGTCTGGCCAGGCAATPAGGACGAGAGGAATTAAGGCTATCAATTAAGGAGAGA 2564

QY 301 GGAAGTCAAAATGTCCTTTGCGATGACATGATTTGATATCTAGAAAACCCCATCT 360

Db 2565 GGAAGTCAAAATGTCCTTTGCGATGACATGATTTGATATCTAGAAAACCCCATCT 2624

QY 361 CTCAGGCCAAATCTCTTATGCTGATAGCAACTTCAGCAAGCTCTAGGATACAAAT 420

Db 2625 CTCAGGCCAAATCTCTTATGCTGATAGCAACTTCAGCAAGCTCTAGGATACAAAT 2684

QY 421 CAATGTACAAAATCACAGCACTCTTATACATCAATTAAGGCTATCAATTAAGGAGAGA 480

Db 2685 CAATGTACAAAATCACAGCACTCTTATACATCAATTAAGGCTATCAATTAAGGAGAGA 2744

QY 481 CATGAGTGAATCTCCCATTCAC 501

Db 2745 CATGAGTGAATCTCCCATTCAC 2765

Search completed: November 7, 2003, 06:15:47  
Job time : 158.728 secs